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(54) Title: GENE CONFERRING DISEASE RESISTANCE IN PLANTS AND USES THEREOF			
(57) Abstract			
<p>The invention concerns the location and characterization of a gene (designated <i>NIM1</i>) which is a key component of the SAR pathway and which in connection with chemical and biological inducers enables induction of SAR gene expression and broad spectrum disease resistance to plants. The invention further concerns plants transformed with the <i>NIM1</i> gene as well as methods employing the gene to create the transgenic plants and employing the gene in a screening assay for compounds capable of inducing broad spectrum disease resistance in plants.</p>			

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GENE CONFERRING DISEASE RESISTANCE IN PLANTS AND USES THEREOF

The present invention relates to disease resistance in plants and to identifying and breeding disease resistance into plants. More particularly, the present invention relates to the identification, isolation and characterization of a gene involved in broad spectrum disease resistance in plants.

Plants are constantly challenged by a wide variety of pathogenic organisms including viruses, bacteria, fungi, and nematodes. Crop plants are particularly vulnerable because they are usually grown as genetically-uniform monocultures; when disease strikes, losses can be severe.

However, most plants have their own innate mechanisms of defense against pathogenic organisms. Natural variation for resistance to plant pathogens has been identified by plant breeders and pathologists and bred into many crop plants. These natural disease resistance genes often provide high levels of resistance to or immunity against pathogens.

In many plant species an initial inoculation by a necrotizing pathogen can immunize the plant to subsequent infection. This acquired disease resistance was first documented in 1901 and is thought to play an important role in the preservation of plants in nature. Particularly well characterized examples of plant immunity are the phenomenon of systemic acquired resistance (SAR) and induced resistance in plants such as tobacco, *Arabidopsis* and cucumber. In these systems, inoculation by a necrotizing pathogen results in systemic protection against subsequent infections by that pathogen as well as a number of other agronomically important bacterial, fungal and viral pathogens.

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Systemic acquired resistance can also be triggered by chemical immunization compounds, certain chemicals that induce the immunity response in plants. Such compounds can be of natural origin, such as salicylic acid (SA), or can be synthetic chemicals, such as 2,6-dichloroisonicotinic acid (INA) and benzo(1,2,3)thiadiazole-7-carbothioic acid S-methyl ester (BTH). Treatment with a pathogen or an immunization compound induces the expression of at least nine sets of genes in tobacco, the best characterized species. Different numbers and types of genes can be expressed in other plants. The level of induction for SAR-related genes induced by immunization compounds is as high as 10,000-fold over background. In particular, SAR is characterized by the expression of SAR genes, including the pathogenesis-related (PR) genes.

The SAR genes are induced following infection by a pathogen. Some of these genes have a role in providing systemic acquired resistance to the plant. These plant proteins are induced in large amounts in response to infection by various pathogens, including viruses, bacteria and fungi. PR proteins were first discovered in tobacco plants (*Nicotiana tabacum*) reacting hypersensitively to infection with tobacco mosaic virus (TMV). Subsequently, PR proteins have been found in many plant species (see Redolfi et al. (1983) Neth J Plant Pathol 89: 245-254; Van Loon (1985) Plant Mol. Biol. 4: 111-116; and Uknes et al. (1992) Plant Cell 4: 645-656.) Such proteins are believed to be a common defensive systemic response of plants to infection by pathogens.

Pathogenesis-related proteins include but are not limited to SAR8.2a and SAR8.2b proteins, the acidic and basic forms of tobacco PR-1a, PR-1b, and PR-1c; PR-1', PR-2, PR-2', PR-2'', PR-N, PR-O, PR-O', PR-4, PR-P, PR-Q, PR-S, and PR-R major proteins; cucumber peroxidases; basic cucumber peroxidase; the chitinase which is a basic counterpart of PR-P or PR-Q; the beta- 1,3-glucanase (glucan endo- 1,3-beta-glucosidase, EC 3.2.1.39) which is a basic counterpart of PR-2, PR-N or PR-O; and the pathogen-inducible chitinase from

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cucumber. Such PR proteins are disclosed, for example, in Uknes et al. (1992) The Plant Cell 4: 645-656 and the references cited therein.

SAR or SAR-like genes are expressed in all plant species exhibiting systemic acquired resistance. Expression of such genes can be determined by probing with known SAR DNA sequences. For example, see Lawton et al. (1992) Proceedings of the Second European Federation of Plant Pathology (1983), In: Mechanisms of Defence Responses in Plants, B. Fritig and M. Legrand (eds), Kluwer Academic Publishers, Dordrecht, pp. 410-420; Uknes et al. (1992) The Plant Cell 4: 645-656; and Ward et al. (1991) The Plant Cell 3: 1085-1094. Methods for hybridization and cloning are well known in the art. See, for example, Molecular Cloning, A Laboratory Manual, 2nd Edition, Vol. 1-3, Sambrook et al. (eds.) Cold Spring Harbor Laboratory Press (1989) and the references cited therein.

Alternatively such SAR or SAR-like genes can be found by other methods such as protein screening, +/- screening, etc. See, for example, Liang and Pardee (1992) Science 257: 967-971; and St. John and Davis (1979) Cell 16: 443.

Despite much research and the use of sophisticated and intensive crop-protection measures, including genetic transformation of plants, losses due to disease remain in the billions of dollars annually. Disease resistance genes have previously been cloned but transgenic plants transformed with these genes would typically be resistant only to a subset of strains of a particular pathogen species. Despite efforts to clone genes involved in SAR, a gene controlling broad spectrum disease resistance has not been isolated and characterized.

Several lines of evidence indicate that endogenously produced SA is involved in the signal transduction pathway coupling the perception of pathogen infection with the onset of SAR. Mutants which retain the ability to accumulate SA in response to pathogen yet have lost the ability to induce SAR genes or resistance after application of SA or INA have been

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described by Delaney, et al., Proc. Natl. Acad. Sci. 92: 6602-6606 (1995) and in WO94/16077 the whole of which is incorporated herein by reference.

It has now been discovered that these mutants contain a mutant gene, which gene in its wildtype form controls SAR gene expression and SAR itself. The present invention recognizes that the mutant gene confers broad spectrum disease susceptibility to mutant plants and renders them noninducible to pathogens and chemical inducers.

The present invention concerns the identification, isolation and characterization of the wildtype (*NIM1*) gene, a gene which allows activation in a plant of SAR and SAR gene expression in response to biological and chemical inducers.

A mutant gene has been identified in Mutagenized *Arabidopsis* plants. These plants have been found to be defective in their normal response to pathogen infection in that they do not express genes associated with systemic acquired resistance (SAR) nor are they capable of exhibiting SAR. These mutants contain a defective gene which has been labelled *nim1* (for noninducible immunity).

The present invention also concerns the use of the cloned *NIM1* gene and variants thereof to create transgenic plants that have broad spectrum disease resistance and to the transgenic plants produced thereby. The invention further concerns the use of the cloned *NIM1* gene and variants thereof in a screening method for identifying compounds capable of inducing broad spectrum disease resistance in plants.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows the effect of chemical inducers on the induction of PR gene expression in wild-type and *nim1* plants.

Figure 2 depicts PR-1 gene expression in pathogen-infected Ws-O and *nim1* plants over the course of 6 days from the initiation of infection.

Figure 3 shows the levels of SA accumulation in Ws-O and *nim1* plants infected with *P. syringae*.

Figure 4 shows the genetic map of the *NIM1* region as determined by AFLP and SSLP analysis.

Figure 5 depicts a physical map of the *NIM1* region as determined by YAC clone analysis.

Figure 6 shows a physical map of an extended P1/BAC contig.

Figure 7 shows a physical map setting forth the positions of P1 and BAC clones with respect to the flanking AFLP markers and YACs.

Figure 8 shows a physical map of a further extended P1/BAC contig containing the *NIM1* gene.

Figure 9 shows an integrated genetic and physical fine map of the *NIM* region.

Figure 10 shows an integrated map of the *NIM1* region.

Figure 11 shows an integrated map of the *NIM1* region including the new AFLP markers.

Figure 12 is a schematic representation of recombinants D169 and C105.

Figure 13 is a global map of the chromosomal region centered on *NIM1* with recombinants indicated, including, BACs, YACs and Cosmids in *NIM1* region.

Figure 14 provides the sequence of the 9.9kb region of clone BAC-04 containing the *NIM1* gene.

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Figure 15 shows the nucleic acid sequence of the *NIM1* gene and the amino acid sequence of the *NIM1* gene product, including changes in the various alleles.

Figure 16 shows the expression of *NIM1* induced by INA, BTH, SA and pathogen in wild type and mutant alleles of *nim1*.

Figure 17 shows the expression of PR-1 in *nim1* mutants and wild-type plants.

Figure 18 shows disease resistance in various *nim1* mutants.

Figure 19 is an amino acid sequence comparison of Expressed Sequence Tag regions of the *NIM1* protein and cDNA protein products of 4 rice gene sequences (see SEQ ID NO: 3).

DEFINITIONS

AA:	Amino Acid
AFLP:	Amplified Fragment Length Polymorphism
avrRpt2:	avirulence gene Rpt2, isolated from <i>Pseudomonas syringae</i>
BAC:	Bacterial Artificial Chromosome
BTH:	benzo(1,2,3)thiadiazole-7-carbothioic acid S-methyl ester
Col:	<i>Arabidopsis</i> ecotype Columbia
ECs:	Enzyme combinations
INA:	2,6-dichloroisonicotinic acid
Ler:	<i>Arabidopsis</i> ecotype <i>Landsberg erecta</i>
<i>NIM1</i> :	the wildtype gene, conferring disease resistance to the plant
<i>nim</i> :	mutant allele of <i>NIM1</i> , conferring disease susceptibility to the plant
<i>nim1</i> :	mutant plant line
ORF:	open reading frame
PCs:	Primer combinations

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SA: salicylic acid
SAR: Systemic Acquired Resistance
SSLP: Simple Sequence Length Polymorphism
Ws-O: *Arabidopsis* ecotype Wassilewskija
YAC: Yeast Artificial Chromosome

The *NIM1* gene has been cloned by mapping and walking techniques which indicate that the gene is contained in a ~105 Kb region. (See Figure 13 and Table 16). This region is delineated by the L84.6b marker on the left and the L84.T2 marker on the right. Only three overlapping cosmids made from wild-type DNA from the 105 Kb region complement the *nim1* mutant phenotype (Figure 13 and Table 16). These three cosmids only overlap in a 9.9 Kb region defined by the left end of cosmid clone D7 and the right end of cosmid D5 as pictured in Figure 13. Many other cosmids made to other areas of the 105 Kb region do not complement the *nim1* phenotype (Figure 13 and Table 16). A near full length cDNA clone to the *NIM1* gene indicates the appropriate intron-exon borders and defines the amino acid sequence of the gene product. Only the *NIM1* gene region within the 9.9 Kb complementing region has sequence changes in various *nim1* mutant alleles (Table 18). Three other potential gene regions showed no sequence changes that are associated with the *nim1* phenotype. The sequence changes found in the *NIM1* gene region are consistent with altered function or loss of function of the gene product. The severity of the change to the *NIM1* gene region in a particular mutant allele is roughly correlated with the observed physiological severity of that *nim1* allele. Only the *NIM1* gene region had detectable RNA (transcription) and this RNA showed abundant changes consistent with the physiological role of *NIM1* in pathogenesis (Table 18 and Figure 16).

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The present invention relates to an isolated gene fragment, the *NIM1* gene, which is a key component of the systemic acquired resistance (SAR) pathway in plants. The *NIM1* gene is associated with the activation of SAR by chemical and biological inducers and, in conjunction with such inducers, is required for SAR and SAR gene expression.

The location of the *NIM* gene is determined by molecular biological analysis of the genome of mutant plants known to carry the mutant *nim1* gene, which gives the host plants extreme sensitivity to a wide variety of pathogens and renders them unable to respond to pathogens and chemical inducers of SAR.

Nim1 mutants are useful as "universal disease susceptible" (UDS) plants by virtue of their being susceptible to many strains and pathotypes of pathogens of the host plant and also to pathogens which do not normally infect the host plant, but which infect other hosts. They can be generated by treating seeds or other biological material with mutagenic agents and then selecting progeny plants for the UDS phenotype by treating progeny plants with known chemical inducers (e.g. INA) of the systemic acquired response and then infecting the plants with a known pathogen. Noninducible mutants develop severe disease symptoms under these circumstances, whereas non-mutants are induced by the chemical compound to systemic acquired resistance. *nim* mutants can be equally selected from mutant populations generated by chemical and irradiation mutagenesis, as well as from populations generated by T-DNA insertion and transposon-induced mutagenesis.

Techniques for generating mutant plant lines are well known in the art. The *nim* plant phenotype is used as a tool to identify an isolated gene fragment which allows expression of broad spectrum disease resistance in plants.

Comprised by the present invention is an isolated DNA molecule comprising a mutant gene of the *NIM1* gene which is a *nim1* gene.

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Following the use of a *nim1* mutant or plant to isolate the wild-type *NIM1* gene necessary for constitutive expression of SAR genes, the resistance trait, in combination with other characteristics important for production and quality, can be incorporated into plant lines through breeding. Breeding approaches and techniques are known in the art. See, for example, Welsh J. R., Fundamentals of Plant Genetics and Breeding, John Wiley & Sons, NY (1981); Crop Breeding, Wood D. R. (Ed.) American Society of Agronomy Madison, Wisconsin (1983); Mayo O., The Theory of Plant Breeding, Second Edition, Clarendon Press, Oxford (1987); Singh, D.P., Breeding for Resistance to Diseases and Insect Pests, Springer-Verlag, NY (1986); and Wricke and Weber, Quantitative Genetics and Selection Plant Breeding, Walter de Gruyter and Co., Berlin 1986).

A further object of the invention is a chimeric gene comprising a promotor active in plant operably linked to a heterologous DNA molecule encoding the aminoacid sequence of a *NIM1* gene product and variants thereof according to the invention.

Methodologies for the construction of plant expression cassettes as well as the introduction of foreign DNA into plants is generally described in the art. Generally, for the introduction of foreign DNA into plants, Ti plasmid vectors have been utilized for the delivery of foreign DNA. Also utilized for such delivery have been direct DNA uptake, liposomes, electroporation, micro-injection, and microprojectiles. Such methods had been published in the art. See, for example, Bilang et al. (1991) Gene 100: 247-250; Scheid et al., (1991) Mol. Gen. Genet. 228: 104-112; Guerche et al., (1987) Plant Science 52: 111-116; Neuhauser et al., (1987) Theor. Appl. Genet. 75: 30-36; Klein et al., (1987) Nature 327: 70-73; Howell et al., (1980) Science 208:1265; Horsch et al., (1985) Science 227: 1229-1231; DeBlock et al., (1989) Plant Physiology 91: 694-701; Methods for Plant Molecular Biology (Weissbach and Weissbach, eds.) Academic Press, Inc. (1988); and Methods in Plant Molecular Biology (Schuler and Zielinski, eds.) Academic Press, Inc. (1989). See also US. patent application

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Serial Nos. 08/438,666 filed May 10, 1995, and WO 93/07278, both of which are incorporated herein by reference in their entirety. It is understood that the method of transformation will depend upon the plant cell to be transformed.

It is further recognized that the components of the expression cassette may be modified to increase expression. For example, truncated sequences, nucleotide substitutions or other modifications may be employed. Plant cells transformed with such modified expression systems, then, would exhibit overexpression or constitutive expression of SAR genes necessary for activation of SAR.

The DNA molecule or gene fragment conferring disease resistance to plants by allowing induction of SAR gene expression can be incorporated in plant or bacterial cells using conventional recombinant DNA technology. Generally, this involves inserting the DNA molecule into an expression system to which the DNA molecule is heterologous (i.e., not normally present). The heterologous DNA molecule is inserted into the expression system or vector in proper orientation and correct reading frame. The vector contains the necessary elements for the transcription and translation of the inserted protein-coding sequences. A large number of vector systems known in the art can be used, such as plasmids, bacteriophage viruses and other modified viruses. Suitable vectors include, but are not limited to, viral vectors such as lambda vector systems Igt11, Igt10 and Charon 4; plasmid vectors such as pBI121, pBR322, pACYC177, pACYC184, pAR series, pKK223-3, pUC8, pUC9, pUC18, pUC19, pLG339, pRK290, pKC37, pKC101, pCDNAII; and other similar systems. The DNA sequences can be cloned into the vector using standard cloning procedures in the art, as described by Maniatis et al., Molecular Cloning: A Laboratory Manual, Cold Spring Laboratory, Cold Spring Harbor, New York (1982).

A further object of the invention is a recombinant vector comprising the chimeric gene according to the invention.

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In order to obtain efficient expression of the gene or gene fragment of the present invention, a promoter must be present in the expression vector. RNA polymerase normally binds to the promoter and initiates transcription of a gene. Promoters vary in their strength, i.e., ability to promote transcription. Depending upon the host cell system utilized, any one of a number of suitable promoters can be used. Suitable promoters include ubiquitin, nos promoter, the small subunit ribulose biphosphate carboxylase gene promoter, the small subunit chlorophyll A/B binding polypeptide promoter, the 35S promoter of cauliflower mosaic virus, and promoters isolated from plant genes. See C.E. Vallejos, et al., "Localization in the Tomato Genome of DNA Restriction Fragments Containing Sequences Homologous to the rRNA (45S), the major chlorophyll _{AB} Binding Polypeptide and the Ribulose Bisphosphate Carboxylase Genes," Genetics 112: 93-105 (1986), which discloses the small subunit materials. The nos promoter and the 35S promoter of cauliflower mosaic virus are well known in the art.

Once the disease resistance gene of the present invention has been cloned into an expression system, it is ready to be transformed into a plant cell. Plant tissues suitable for transformation include leaf tissues, root tissues, meristems, and protoplasts.

Bacteria from the genus *Agrobacterium* can be utilized to transform plant cells. Suitable species of such bacterium include *Agrobacterium tumefaciens* and *Agrobacterium rhizogens*. *Agrobacterium tumefaciens* (e.g., strains LBA4404 or EHA105) is particularly useful due to its well-known ability to transform plants.

Another approach to transforming plant cells with a gene involves propelling inert or biologically active particles at plant tissues and cells. This technique is disclosed in U.S. Patent Nos. 4,945,050; 5,036,006; and 5,100,792 all to Sanford et al. Generally, this procedure involves propelling inert or biologically active particles at the cells under conditions effective to penetrate the outer surface of the cell and afford incorporation within

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the interior thereof. When inert particles are utilized, the vector can be introduced into the cell by coating the particles with the vector containing the desired gene. Alternatively, the target cell can be surrounded by the vector so that the vector is carried into the cell by the wake of the particle. Biologically active particles (e.g., dried yeast cells, dried bacterium or a bacteriophage, each containing DNA sought to be introduced) can also be propelled into plant cell tissue.

The isolated gene fragment of the present invention can be utilized to confer disease resistance to a wide variety of plant cells, including those of gymnosperms, monocots, and dicots. Although the gene can be inserted into any plant cell falling within these broad classes, it is particularly useful in crop plant cells, such as rice, wheat, barley, rye, corn, potato, carrot, sweet potato, sugar beet, bean, pea, chicory, lettuce, cabbage, cauliflower, broccoli, turnip, radish, spinach, asparagus, onion, garlic, eggplant, pepper, celery, carrot, squash, pumpkin, zucchini, cucumber, apple, pear, quince, melon, plum, cherry, peach, nectarine, apricot, strawberry, grape, raspberry, blackberry, pineapple, avocado, papaya, mango, banana, soybean, tobacco, tomato, sorghum and sugarcane.

The expression system of the present invention can be used to transform virtually any crop plant cell under suitable conditions. Transformed cells can be regenerated into whole plants such that the gene imparts disease resistance to the intact transgenic plants. As set forth above, the expression system can be modified so that the disease resistance gene is continuously or constitutively expressed.

Transformation

The present system can be utilized in any plant which can be transformed and regenerated. Such methods for transformation and regeneration are well known in the art. As well as the above cited references, see also, An, G., Watson, B.D., and Chiang, C.C.

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As *nim1* host plants may also be susceptible to pathogens outside of the host-range into which they normally fall, these plants also have significant utility in the molecular, genetic, and biological study of host-pathogen interactions. Furthermore, the UDS phenotype of *nim1* plants also renders them of utility for fungicide screening. *nim1* mutants selected in a particular host have considerable utility for the screening of fungicides using that host and pathogens of the host. The advantage lies in the UDS phenotype of the mutant, which circumvents the problems encountered due to hosts being differentially

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susceptible to different pathogens and pathotypes, or even resistant to some pathogens or pathotypes.

Pathogens of the invention include but are not limited to viruses or viroids, e.g. tobacco or cucumber mosaic virus, ringspot virus or necrosis virus, pelargonium leaf curl virus, red clover mottle virus, tomato bushy stunt virus, and like viruses; fungi, e.g. *Phytophthora parasitica* and *Peronospora tabacina*; bacteria, e.g. *Pseudomonas syringae* and *Pseudomonas tabaci*; insects such as aphids, e.g. *Myzus persicae*; and lepidoptera, e.g., *Heliothus spp.*; and nematodes, e.g., *Meloidogyne incognita*. The methods of the invention are useful against a number of disease organisms of maize including but not limited to downy mildews such as *Sclerophthora macrospora*, *Sclerophthora rayissiae*, *Sclerospora graminicola*, *Peronosclerospora sorghi*, *Peronosclerospora philippinensis*, *Peronosclerospora sacchari* and *Peronosclerospora maydis*; rusts such as *Puccinia sorphi*, *Puccinia polysora* and *Physopella zeae*; other fungi such as *Cercospora zeae-maydis*, *Colletotrichum graminicola*, *Fusarium moniliforme*, *Gibberella zeae*, *Exserohilum turcicum*, *Kabatiellu zeae* and *Bipolaris maydis*; and bacteria such as *Erwinia stewartii*.

DESCRIPTION OF THE SEQUENCE LISTING

SEQ ID NO:1 - 9919-bp genomic sequence from Figure 14.

SEQ ID NO:2 - 5655-bp genomic sequence from Figure 15.

SEQ ID NO:3 - AA sequence of wild-type NIM protein encoded by cds of seq id no:2.

SEQ ID NO:4 - Rice-1 AA sequence 33-155 from Figure 19.

SEQ ID NO:5 - Rice-1 AA sequence 215-328 from Figure 19.

SEQ ID NO:6 - Rice-2 AA sequence 33-155 from Figure 19.

SEQ ID NO:7 - Rice-2 AA sequence 208-288 from Figure 19.

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SEQ ID NO:8 - Ric -3 AA sequence 33-155 from Figur 19.

SEQ ID NO:9 - Rice-3 AA sequence 208-288 from Figure 19.

SEQ ID NO:10 - Rice-4 AA sequence 33-155 from Figure 19.

SEQ ID NO:11 - Rice-4 AA sequence 215-271 from Figure 19.

DEPOSITS

The following vector molecules have been deposited with American Type Culture Collection 12301 Parklawn Drive Rockville, MD 20852 U.S.A. on the dates indicated below:

Plasmid BAC-04 was deposited with ATCC on May 8, 1996 as ATCC 97543.

Plasmid P1-18 was deposited with ATCC on June 13, 1996 as ATCC 97606.

Cosmid D7 was deposited with ATCC on September 25, 1996 as ATC 97736.

EXAMPLES

Example 1

Identifying *NIM1* clones by map-based cloning. High resolution genetic mapping and physical mapping of *NIM1* in Arabidopsis.

1. Plant Material and Isolation of *nim1* Mutants.

Nim1 mutants were isolated from two Arabidopsis ecotype Ws-O plant populations, as described by Delaney et al., (1995) PNAS 92, 6602-6606. One mutant population was in the form of an M2 library derived from ethyl methane sulfonate (EMS) mutagenized seeds (purchased from Lehle, Round Rock, TX) and the other was in the form of a T-DNA population derived from seed obtained from the Ohio State University Arabidopsis Biological Resource Center (Columbus, OH).

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The basis of the screen for noninducible immunity (*nim1*) mutants was to screen mutagenized plant populations for plants in which resistance to a virulent pathogen could not be induced by INA (2,6 dichloro isonicotinic acid; Metraux, et al., 1991. In: Advances in Molecular Genetics of Plant-Microbe Interactions. Vol 1, 432-439. Hennecke and Verma, eds.; Kessmann et al. 1993 In: Mode of action of agrochemicals. Y Honma, ed.; Vernooij et al, 1995, Molec. Pl. Microbe Interaction 8, 228-234).

Plants from the mutant populations were grown at high density in large trays in commercial planting mix. When the plants were 2 weeks of age, the trays were sprayed with 0.25 mg/ml INA. Four days later, the plants were sprayed with a spore suspension of *Peronospora parasitica*, isolate EmWa (EmWa), at 5×10^4 to 1×10^5 spores/ml. This fungus is normally virulent on the Arabidopsis Ws-O ecotype, unless resistance is first induced in these plants with INA or a similar compound.

Following incubation in a high humidity environment, plants with visible disease symptoms were identified, typically 7 days after the infection. These plants did not show resistance to the fungus, despite the application of the resistance-inducing chemical and were thus potential *nim* (non-immunity) mutant plants. From 360,000 plants, 75 potential *nim* mutants were identified.

These potential mutant plants were isolated from the flat, placed under low humidity conditions and allowed to set seed. Plants derived from this seed were screened in an identical manner for susceptibility to the fungus EmWa, again after pretreatment with INA. The progeny plants that showed infection symptoms were defined as *nim* mutants. Six *nim* lines were thus identified. One line (*nim1*) was isolated from the T-DNA population and five from the EMS population.

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2. Scoring Plant Reactions to INA and Other Chemical Inducers of Disease Resistance.

i. Phenotypic analysis of *nim1*.

Salicylic acid (SA) and benzo(1,2,3)thiadiazole-7-carbothioic acid S-methyl ester (BTH) are two chemicals that, like INA, induce broad spectrum disease resistance, termed Systemic Acquired Resistance (SAR), in wildtype plants. Since INA did not induce resistance in the *nim1* plants, these plants were also evaluated for their disease resistance response following pretreatment with SA and BTH (as partly described in Delaney et al, 1995, PNAS 92, 6602-6606).

Plants were sprayed with 1, 5, or 15 mM SA or 0.25 mg/ml BTH and challenge inoculated with EmWa 5 days later (as described in example 1 above). Both SA and BTH failed to protect *nim1* plants from fungal infection, as evidenced by the presence of disease symptoms and fungal growth on these plants. Thus, the *nim1* plants were not responsive to any of the SAR-inducing chemicals, implying that the mutation was downstream of the entry point(s) for these chemicals in the resistance induction pathway.

Nim1 was also evaluated for its disease response to infection with 2 incompatible *P. parasitica* isolates, Wela and Noco (i.e. these fungal strains do not cause disease on wildtype Ws-O plants). *nim1* plants were sprayed with conidial suspensions of $5-10 \times 10^4$ spores/ml of Wela or Noco and incubated under high humidity for 7 days. Unlike wildtype plants, *nim1* plants developed disease symptoms in response to both Wela and Noco infection. The symptoms were necrotic flecking and trailing, with some sporulation. Following lactophenol blue staining, fungal hyphae were easily observed in the leaves of *nim1* plants. Thus, the *nim1* plants are susceptible to normally incompatible *P. parasitica* isolates. This result shows that the *nim1* plants are not only defective in chemically induced

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disease resistance, but are also defective in natural resistance to microorganisms that are normally not pathogenic.

ii. Biochemical analysis of *nim1*.

SA, INA and BTH induce SAR and expression of the SAR genes, which include the Pathogenesis Related genes PR-1, PR-2 and PR-5 in *Arabidopsis*. Since these compounds did not induce disease resistance in *nim1* (as described in example 1.2 above), this mutant line was analyzed for SAR gene expression following SA, INA or BTH treatment.

After treatment of *nim1* plants with SA, INA or BTH, plant tissue was harvested and analyzed for accumulation of RNA from the PR-1, PR-2 and PR-5 genes. To this end, total RNA was isolated from the treated tissues and electrophoresed on an agarose gel. Triplicate gel blots were prepared and each was hybridized with a probe for one of these 3 SAR genes as described in Delaney et al, 1995, PNAS 92, 6602-6606. In contrast to the case in wildtype plants, the chemicals did not induce RNA accumulation from any of these 3 SAR genes in *nim1* plants, as shown in figure 1. Taken together, the results indicate that the chemicals induce neither SAR nor SAR gene expression in *nim1* plants.

Since the chemicals did not induce SAR, or SAR gene expression in *nim1* plants, it was of interest to investigate whether pathogen infection could induce SAR gene expression in these plants, as it does in wildtype plants. Ws-O and *nim1* plants were sprayed with EmWa spores as described and tissue collected for RNA analysis at several timepoints. Pathogen infection (EmWa) of wildtype Ws-O plants induced PR-1 gene expression within 4 days after infection, as shown in figure 2. In *nim1* plants, however, PR-1 gene expression is not induced until 6 days after infection and the level is reduced relative to the wildtype at that time. Thus, following pathogen infection, PR-1 gene expression in *nim1* plants is delayed and reduced relative to the wildtype.

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Infection of wildtype plants with pathogens that cause a necrotic reaction leads to accumulation of SA in the infected tissues. It has been shown that this endogenous SA is required for signal transduction in the SAR pathway, i.e., breakdown of the endogenous SA leads to a decrease in disease resistance. This defines SA accumulation as a marker in the SAR pathway (Gaffney et al, 1993, *Science* 261, 754-756).

Nim1 plants were tested for their ability to accumulate SA following pathogen infection. *Pseudomonas syringae* tomato strain DC 3000, carrying the *avrRpt2* gene, was injected into leaves of 4-week-old *nim1* plants. The leaves were harvested 2 days later for SA analysis as described by Delaney et al, 1995, *PNAS* 92, 6602-6606. This analysis showed that the *nim1* plants accumulated high levels of SA in infected leaves, as shown in Figure 3. Uninfected leaves also accumulated SA, but not to the same levels as the infected leaves, similar to what has been observed in wild-type *Arabidopsis*. This indicated that the *nim* mutation maps downstream of the SA marker in the signal transduction pathway. This was anticipated, as INA and BTH (inactive in *nim1* plants) are known to stimulate a component in the SAR pathway downstream of SA (Vernooij et al., 1995, *Molec. Pl. Microbe Interaction* 8, 228-234; Friedrich, et al., 1996, *The Plant Journal* 9, in press; and Lawton, et al., 1996, *The Plant Journal* 9, in press). In addition, as described in Example 1.2, exogenously applied SA did not protect *nim1* from EmWa infection.

3. Genetic analysis of *nim1*.

Nim1 plants were backcrossed to wild-type Ws-O plants, and F1 progeny were tested for resistance to EmWa after INA pretreatment, as described in Example 1.1 above. None of the INA-pretreated F1 plants had infection symptoms, whereas the *nim1* control plants did show infection. Hence, the *nim1* mutation was determined to be recessive.

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The F₂ population from the Ws-O x *nim1* cross was also assayed for its disease resistance after INA pretreatment. Of this population, approximately 1/4 (32/130 plants) showed disease symptoms after EmWa treatment of INA pre-sprayed plants and 3/4 (98/130 plants) showed no disease. These results indicate that the *nim* mutation identifies a single genetic locus and corroborates the F₁ data that show the recessive nature of the mutation.

4. Identification of markers in and genetic mapping of the *NIM* locus.

For conventional map-based cloning of the *NIM* gene, markers had to be identified that were genetically closely linked to the mutation. This was accomplished in 2 steps. First, the *nim1* plants were crossed to a different Arabidopsis genotype, Landsberg erecta (Ler), and F₂ plants from this cross which had a *nim1* phenotype (i.e. plants that are homozygous *nim/nim* at the *NIM* locus) were identified. From these, plants that had a Ler genotype at a nearby DNA marker were identified by molecular analysis. These plants, by virtue of the identification criterion, are recombinant between the marker and the *NIM* locus. The frequency of recombinants defines the genetic distance between the marker and the *NIM* locus.

The second prerequisite for map based cloning is that markers are identified that are genetically very close to the *NIM* locus, i.e. markers that identify very few recombinants. If genetic markers are identified that are very close, then these can be used to isolate genomic DNA clones that are close to the *NIM* locus. The *NIM* locus can then be cloned by walking, if not already present on the cloned DNA. Walking can be initiated from both sides of the gene. It relies on obtaining overlapping clones that are successively closer to the gene of interest. When a single DNA marker is obtained from a walk initiated from, for instance, the North end and it identifies no recombinants between this marker and the gene of interest, it must be very close to the gene. However, if the marker does identify

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recombinant(s) from the South end, the clone from which the marker was obtained must have crossed the gene. By definition then, the gene of interest is cloned. It must be located between this marker and the last North-end marker that identifies the least number of recombinants from the North end.

In a first step, a large number of recombinants are generated by genetic crossing. In a second step, recombinants that are close to the *NIM* gene are identified with the use of molecular markers. Many markers have been described in the literature and several methods exist to develop additional markers. Our approach has relied on a number of marker systems, including SSLPs and AFLPs (see below).

i. Genetic crosses.

In order to map the chromosomal position of the *NIM* gene relative to the SSLP and AFLP markers, *nim1* was crossed to Ler to make a mapping population. F₂ plants from this cross were grown and leaves harvested for future DNA extractions. Next, the F₂ plants were scored for the *nim1* phenotype, as described in example 1.1 above. Also, F₃ populations derived from individual F₂ plants were grown and scored for the *nim* phenotype. DNA was extracted from the stored tissue of *nim1* phenotype F₂ and F₃ plants by the CTab method, as described (Rogers and Bendich, 1988, Plant Molecular Biology Manual, A6, 1-10). This DNA was used for mapping the *NIM* gene, as described below.

ii. Simple Sequence Length Polymorphism markers.

The Simple Sequence Length Polymorphism (SSLP) markers ATHGENEA and nga111 have been described (Bell and Ecker, 1994, Genomics 19, 137-144). Primers used for detection of these SSLPs are listed in Table 1.

Table 1. SSLP primer sequences.

primer set	primer sequence (5' to 3')
ATHGENEA (1)	ACC ATG CAT AGC TTA AAC TTC TTG ACA TAA CCA CAA ATA GGG GTG C
ATHGENEA (2)	ACC ATG CAT AGC TTA AAC TTC TTG CCA AAT GTC AAA ATA CTC GTC
nga111 (1)	CTC CAG TTG GAA GCT AAA GGG TGT TTT TTA GGA CAA ATG GCG
nga111 (2)	CTC CAG TTG GAA GCT AAA G TGT TTT TTA GGA CAA ATG G

Genetic mapping of the *NIM* gene relative to marker ATHGENEA.

Using the ATHGENEA (1) primers for PCR amplification of Ler genomic DNA, a 205-basepair (bp) band was expected, whereas with Ws-O genomic DNA a band of 211 bp was expected (Bell and Ecker, 1994, Genomics 19, 137-144). The amplification products proved to be difficult to separate on conventional agarose gels. Hence, two alternative methods were developed for separation and detection of these PCR fragments.

In a first method, primerset ATHGENEA (1) (Table 1) was used to amplify genomic DNA in the presence of 6-carboxyrhodamine-labelled UTP (dUTP-R110, obtained from ABI), yielding rhodamine-labelled PCR fragments. The PCR reactions were analyzed on a DNA Sequencer, which detects DNA fragments with single nucleotide resolution.

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The specific reagents were: 1xPCR buffer, 2 mM MgCl₂, dNTPs each 200 mM, 2 mM dUTP-R110, ATHGENEA (1) primers at 0.75 mM, 10 ng DNA and 0.75 units Taq polymerase in a 20 ml reaction volume. Amplification conditions were: 3 minutes 94°C followed by 35 cycles of 15 seconds at 94°C, 15 seconds at 55°C and 30 seconds at 72°C. These samples were analyzed on an ABI 377 DNA Sequencer, capable of detecting fluorescently labeled DNA fragments with single nucleotide (nt.) resolution. This allowed for genotyping the plant samples: a 205-nucleotide DNA fragment was obtained from Ler DNA and a 211-nucleotide band from Ws-O DNA. Thus, DNA fragments differing by 6 nucleotides in length could be easily distinguished, allowing for easy genotyping of samples as homozygous Ws-O, homozygous Ler and heterozygous Ws-O/Ler at the ATHGENEA locus.

In order to increase the throughput of this system, a multiplexing scheme was used. Some DNA samples were PCR amplified as described above with primer set ATHGENEA (1), whereas other samples were analyzed with primerset ATHGENEA (2) (listed in table 2), in each case in the presence of 6-carboxyrhodamine-labeled dUTP. Primer set ATHGENEA (2) was made based on the published sequence of ATHGENEA (Simoens et al., 1988, Gene 67, 1-11). This primerset amplified a DNA fragment of 139 bp from Ler DNA and a 145-bp band from Ws-O DNA. Amplification reaction conditions for primerset ATHGENEA (2) were identical to those described for primerset ATHGENEA (1), above.

Single reactions using primer set ATHGENEA (1) and single reactions using primer set ATHGENEA (2) were mixed together before electrophoresis on the ABI 377 DNA Sequencer. This multiplexing approach allowed for genotyping 2 samples in a single lane of the Sequencer, one at positions 145/139 nt. and one at positions 211/205 nt. on the Sequencer.

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In the second method, PCR fragments were labelled by using a primer labelled with the fluorescent dye FAM-6 (6-carboxyfluorescein) (Integrated DNA Technologies, Inc.). The forward ATHGENEA primers of the ATHGENEA (1) and (2) primer sets are identical in sequence (see Table 1). This primer was labeled with FAM-6 and used in a PCR amplification reaction with the following reagents (Perkin Elmer): 1xXL buffer, 1 mM MgCl₂, dNTPs each at 200 mM, primers each at 0.50 mM (forward primer FAM-6 labeled), 10 ng genomic DNA and 0.5 units XL polymerase in a 20-ml reaction volume. The cycling conditions were: 3 minutes at 94°C, followed by 35 cycles of 15 seconds at 94°C, 15 seconds at 59°C and 30 seconds at 72°C. Again, single reactions using primer set ATHGENEA (1) and single reactions using primer set ATHGENEA (2) were mixed together before electrophoresis on the ABI 377 DNA Sequencer. This multiplexing approach allowed for genotyping 2 samples in a single lane of the Sequencer, one at positions 145/139 nt. and one at positions 211/205 nt.

All F2 and F3 samples from *nim1* phenotype plants were scored for their genotype at the ATHGENEA locus as described above. All samples that were heterozygous at this locus identified plants that were recombinant between the *NIM1* locus and the ATHGENEA locus. In a population of 1144 F2 *nim1* phenotype plants and F3 *nim1* phenotype populations that were scored in this way, 98 were heterozygous at the ATHGENEA locus, giving an estimate of the genetic distance between this SSLP locus and the *NIM1* locus of 4.3 cM. This established that the *NIM1* locus was on chromosome 1, near the ATHGENEA marker.

Genetic mapping of the *NIM1* gene relative to marker nga111.

Two primer sets for SSLP marker nga111 (described in Bell and Ecker, 1994, Genomics 19, 137-144) were used to amplify genomic DNA of F2 and F3 *nim1* phenotype

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plants and control Ws-O and Ler plants. Primer set nga111 (1) (described in Bell and Ecker, 1994, *Genomics* 19, 137-144 and listed in Table 1) was used under the following conditions: 1xPCR buffer, 2 mM MgCl₂, dNTPs each at 200 mM, primers at 0.75 mM, 10 ng DNA and 0.75 unit Taq polymerase in a 20 ml reaction volume. Primer set nga111 (2) (listed in Table 1, and a derivative of primer set nga111 (1)) was used under different conditions: 1xPCR buffer, 1.5 mM MgCl₂, dNTPs each 200 mM, primers at 1 mM, 10 ng DNA and 1 unit Taq polymerase in a 20 ml reaction volume. Both reactions were amplified by incubation at 94°C for 1 minute, followed by 40 cycles of 15 seconds at 94°C, 15 seconds at 55°C and 30 seconds at 72°C.

The samples were analyzed on 3-5% agarose gels. The band obtained from amplification of Ws-O DNA with either primer set was 146 bp, whereas amplifying Ler DNA resulted in a 162-bp band. Plant samples that were heterozygous at the nga111 locus identified plants that were recombinant between this SSLP marker and the *NIM* locus. Among 1144 F2 *nim1* phenotype plants and F3 *nim1* phenotype populations, 239 were identified as heterozygous for the nga111 marker, giving an estimate for the genetic distance between the SSLP marker and the *NIM* locus of 10.4 cM. This corroborated that the *NIM1* locus was on chromosome 1. Since few *nim1* phenotype plants existed that were heterozygous at both ATHGENEA and nga111, the *NIM1* locus was determined to be between these 2 markers, with ATHGENEA located North of the *NIM1* gene and nga111 located South of the *NIM1* gene. This placed the *NIM1* gene approximately 10 cM north of nga111, near position 85 on chromosome 1 (Lister and Dean, 1993, *Plant J.* 4, 745-750; Bell and Ecker, 1994, *Genomics* 19, 137-144).

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iii. Amplified Fragment Length Polymorphism markers.

For map-based cloning of the *NIM1* gene, it is necessary to identify molecular markers that are successively closer to this gene. For this purpose, Amplified Fragment Length Polymorphism (AFLP) markers were generated by using the selective restriction fragment amplification method described by Zabeau and Vos (1993, European Patent Application EP 0534858) and Vos et al. (1995, Nucleic Acid Research 23, 4407-4414).

Outline of the AFLP Technology.

The use of the AFLP technology in mapping relies on selective amplification of a set of DNA bands in 2 genetically distinct samples. Finding that any of the obtained bands are different between the 2 genotypes identifies those bands as markers for that genotype. If the marker cosegregates at high frequency with the gene (mutation) of interest, then the marker is close to the genetic locus.

Selective amplification of a small set of DNA fragments in a complex DNA sample is achieved in a 2-step process. First, DNA fragments are generated by digesting the DNA with restriction enzymes, followed by ligation of adapters to the ends. Second, primers consisting of a sequence complementary to the adapters plus a 3' extension (typically 0-3 nucleotides) are used to amplify only those DNA fragments with ends that are complementary to these primers. If a single nucleotide extension is used, then theoretically, each primer will "fit" on approximately 1/4 of all fragments, with 1/16 of all fragments having a primer fit on both ends. Thus, a limited set of DNA fragments is amplified with these primers. By further radiolabelling one primer, an even smaller subset of visible bands can be obtained.

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AFLP analysis.

For AFLP analysis of DNA samples, 50 ng DNA was digested with the appropriate enzymes (usually EcoRI and MseI; see below) and adapters (listed in table 2 below) were ligated to the restriction fragments (usually EcoRI and MseI). The sequences of the primers and the YAC, P1 and BAC clones are described in detail below. The templates were used for amplification reactions (approximately 0.5 ng DNA per reaction), using primers that were complementary to the adapters, with short 3' extensions (2 or 3 nucleotides; primer sequences are listed below). Since one of the primers is radioactively labelled (usually the EcoRI primer), only a subset of the amplified fragments is visible upon autoradiographic analysis of the gel used to separate the bands.

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Amplification conditions for cloned DNA (YAC, P1, cosmid) were as follows: 36 cycles of 30 sec. at 94°C (denaturation), 30 sec annealing and 1 min extension at 72°C. The annealing temperature in the first cycle was 65°C and was reduced by 0.7°C in each cycle for the next 12 cycles and then kept at 56°C. For genomic DNA of *Arabidopsis* plants, the amplification was performed in 2 steps: in the first step (preamplification), the DNA was amplified with primers that have a single nucleotide extension (neither primer was labeled). Reaction conditions for this amplification reaction were: 20 cycles of 30 sec. denaturation (94°C), 1 min annealing (56°C) and 1 min extension (72°C). In the second step, the first amplification reaction was diluted 10 fold and reamplified 36 cycles with primers containing the full-size extensions (using one labeled primer) under the following conditions: 30 sec. at 94°C (denaturation), 30 sec annealing and 1 min extension at 72°C. The annealing temperature in the first cycle was 65°C and was reduced by 0.7°C in each cycle for the next 12 cycles and then kept at 56°C. The final reaction products were separated on a polyacrylamide gel and the gel was exposed to film, allowing visualization of the radiolabeled PCR bands. When this procedure was applied to DNA from 2 genotypes simultaneously, AFLP bands were identified that were diagnostic for one genotype or the other. Such bands are called informative AFLP bands, or AFLP markers. Table 2 shows Adapters used in the AFLP analysis.

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Table 2

enzyme	adapter
EcoRI	5'-CTCGTAGACTGCGTACC-3' 3'-CATCTGACGCATGGTTAA-5'
HindIII	5'-CTCGTAGACTGCGTACC-3' 3'-CATCTGACGCATGGTCGA-5'
PstI	5'-CTCGTAGACTGCGTACATGCA-3' 3'-CATCTGACGCATGT-5'
MseI	5'-GACGATGAGTCCTGAG-3' 3'-TACTCAGGACTCAT-5'

Generation of AFLP markers and fine mapping of the *NIM1* locus.

A population of recombinant inbred lines derived from a cross between the *Arabidopsis* ecotypes Landsberg erecta (Ler) and Columbia (Col) (Lister and Dean, 1993, Plant J. 4, 745-750) was used for AFLP marker screening. The primers used for the AFLP screening were:

EcoRI-primers: 5'-GACTGCGTACCAATTCWN-3'

MseI-primers: 5'-GATGAGTCCTGAGTAAXWN-3'

An "N" in the primers indicates that this part was variable (A, C, G or T), a "W" indicates A or T, and an "X" indicates a C. All 8 possible primers were used for both the

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EcoRI- and MseI-primer. This gave a total of 64 (8 x 8) primer combinations (PCs) that were used to amplify DNA from the recombinant inbred line and the parental genotypes, Ler and Col, as described above. The amplification reactions were run on a denaturing polyacrylamide gel to separate AFLP fragments by size and the gel was exposed to film. The film was inspected for bands that were present in only one genotype, i.e. inspected for AFLP markers.

The AFLP markers, i.e., DNA fragments that are polymorphic between both parents of the recombinant inbred lines, were used for constructing a genetic map of the recombinant inbred line population. Example 1.5i below describes the mapping of the *NIM1* gene on *Arabidopsis* chromosome 1, at approximately position 85. Those AFLP markers that had been mapped (using the recombinant inbred line) between positions 81 and 88 of *Arabidopsis* chromosome 1 were chosen for analyzing recombinant plants for the presence of said AFLP markers and thus for mapping the *NIM1* gene more precisely. Seven AFLP markers from this region were identified as being informative; they were polymorphic between both parents of the *nim1*xLer cross. Six AFLP markers were Ler-specific, i.e. these AFLP markers were absent in Ws (and in Col as well). One AFLP marker was Ws-specific, i.e. a Col-specific AFLP marker (absent in Ler) was also present in Ws. These AFLP markers are: L81.1, L81.2, W83.1, L84, L85, L87 and L88 (an L-marker is specific for ecotype Ler and a W-marker is specific for both the ecotypes Col and Ws; the number indicates the map position). These AFLP markers were used to analyze recombinant plants from the *nim1*xLer cross (see below). In addition, AFLP marker C86 (a recombinant, inbred-line-derived marker specific for Col) was used in isolating DNA clones (see below). Table 3 lists the primer sequences that were used to obtain these AFLP markers.

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Table 3 shows primer combinations of AFLP markers derived from recombinant inbred line population.

"EcoRI-" refers to the sequence 5'-GACTGCGTACCAATTC-3' and

"MseI-" refers to the sequence 5'-GATGAGTCCTGAGTAA-3'.

Table 3

AFLP marker	Corresponding primer combinations	
L81.1	EcoRI-CA	MseI-CCG
L81.2	EcoRI-AA	MseI-CAA
W83.1	EcoRI-CA	MseI-CTC
L84	EcoRI-AAT	MseI-CAA
L85	EcoRI-CA	MseI-CCT
L87	EcoRI-CA	MseI-CTT
L88	EcoRI-AG	MseI-CTA
C86	EcoRI-AG	MseI-CCT

A detailed genetic map of the region was constructed using the AFLP markers described above by typing the recombinants. A total of 337 recombinant plants were available out of 1144 F2 *nim1* plants. These recombinants were first screened with the North-flanking AFLP markers L81.2 and ATHGENEA and the South-flanking markers L88 and *nga111*. Forty-eight plants were homozygous *nim1/nim1* and heterozygous at ATHGENEA and L81.2, and 21 plants were homozygous *nim1/nim1* and heterozygous at *nga111* and L88. These recombinant plants were further analyzed with 9 AFLP markers in the *NIM* region, including 4 AFLP markers that were derived from the recombinant inbred line mapping population, (W83.1, L84, L85 and L87) and 5 AFLP markers derived from analysis of YAC clones (W83.3/W84.1, W84.2, W85.1, W86.1 and L86, see below).

The genetic map of *NIM1*, based on this analysis, is depicted in Figure 4. As seen, 27 recombinants were found between marker W84.2 and *NIM1* and 14 recombinants were

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found between W85.1 and *NIM1*. Marker L85 is linked closely to *NIM1*, but this marker could not be mapped on the YAC, BAC or P1 clones (see below) and was, therefore, not useable for identification of the *NIM1* gene.

5. Physical mapping of the *NIM1* region.

i. Isolation of YAC clones using AFLP markers closely linked to *NIM1*.

The CIC library, an Arabidopsis ecotype Columbia YAC library (Bouchez et al, 1995, 6th Int. Conf on Arabidopsis Research, Madison, WI), was screened for YAC clones in the *NIM* region. This library has about 2.5 nuclear genome equivalents and has an average insert size of 450 kb. The YAC library was screened with two AFLP markers: W83.1 and C86. W83.1 is the most closely linked recombinant, inbred-line-derived AFLP marker north of *NIM1*, and C86 is a recombinant, inbred-line-derived AFLP marker specific for Col (absent in Ler and Ws). C86 mapped south of the *NIM1* gene on the map of the recombinant inbred line population. This Col AFLP marker has been used instead of the closely linked Ler AFLP markers (Figure 4), because the latter AFLP markers detected only ecotype *Landsberg erecta* and hence cannot be used for screening the Columbia YAC library.

The YAC library was screened in two steps. Firstly, the cells of the YAC clones of each plate of the twelve 96-well microtiter plates were pooled (a plate pool) and used for DNA isolation as described by Ross et al (1991, Nucleic Acids Res. 19, 6053). The pools were screened with both AFLP markers. Subsequently, from each positive plate pool, the DNA samples of each row (a pool of 8 clones) and of each column (a pool of 12 clones) were screened with the AFLP marker for which the plate pool was positive. In this way, the individual positive YAC clones could be identified. The screening yielded a total of 4 YAC clones: YAC 12F04 and YAC 12H07 were isolated using the North AFLP marker W83.1,

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and YAC 10G07 and YAC 7E03 using the south AFLP marker C86 (for the nomenclature of the YAC clones the CIC numbering is used). The YACs were "fingerprinted" by AFLP, giving YAC-specific AFLP fragments. Fingerprints of the YACs were compared and used to estimate overlaps between the YACs (see also Tables 5 and 6). Based on the AFLP fingerprints, clone 7E03 is essentially covered by clone 10G07 (see also table 5) and clone 12H07 is likewise essentially covered by clone 12F04 (see also table 6).

ii. Generation of AFLP markers from YAC clones.

Since the AFLP markers described above were genetically relatively far from the *NIM1* gene (see figure 3), additional AFLP markers were developed in an effort to find markers that were closer to the *NIM* gene.

A screening for additional YAC-derived AFLP markers was performed on DNA samples of the following: DNA of the isolated YAC clones (4 YACs were identified, as described above), the yeast strain without a YAC, and the three Arabidopsis ecotypes Col, Ler and Ws. In this way, the fragments specific for the YAC clones (absent in the yeast strain and present in Col) could be tested for polymorphism in Ler and Ws (the parents of the recombinant plants identified in Example 1.5 below). All identified polymorphic fragments would thus be additional AFLP markers. In the first AFLP screening the enzyme combination (EC) EcoRI/MseI was used. In this screening two YAC clones, 10G07 and 7E03 (detected with AFLP marker C86, see below), the yeast strain without a YAC and the three Arabidopsis ecotypes Col, Ler and Ws were assayed. The primer combinations with the selective extensions used can be divided into three groups and are depicted in Table 4. A total of 256 (64 + 96 + 96) primer combinations were screened.

In Table 4 below the primer sequences used in the AFLP screening of two YAC clones, 10G07 and 7E03, the yeast strain without a YAC, and the three Arabidopsis

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ecotypes Col, Ler and Ws are shown. Three groups of primer combinations have been used. An "N" in the primers indicates that this part was variable (A, C, G or T), an "S" indicates C or G, a "W" indicates A or T, and a "Y" indicates C or T.

Table 4

EcoRI-primers:

5'-GACTGCGTACCAATTCGW-3'

5'-GACTGCGTACCAATTCTS-3'

MseI-primers:

5'-GATGAGTCCTGAGTAAAAS-3'

5'-GATGAGTCCTGAGTAAASA-3'

5'-GATGAGTCCTGAGTAAATN-3'

5'-GATGAGTCCTGAGTAACAN-3'

5'-GATGAGTCCTGAGTAACTN-3'

EcoRI-primers:

5'-GACTGCGTACCAATTCAN-3'

5'-GACTGCGTACCAATTCCW-3'

5'-GACTGCGTACCAATTCTW-3'

MseI-primers:

5'-GATGAGTCCTGAGTAAAAS-3'

5'-GATGAGTCCTGAGTAAASA-3'

5'-GATGAGTCCTGAGTAAGAY-3'

5'-GATGAGTCCTGAGTAAGTW-3'

5'-GATGAGTCCTGAGTAATCG-3'

5'-GATGAGTCCTGAGTAATCT-3'

5'-GATGAGTCCTGAGTAATGW-3'

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EcoRI-primers:

5'-GACTGCGTACCAATTCGW-3'

5'-GACTGCGTACCAATTCTN-3'

MseI-primers:

5'-GATGAGTCCTGAGTAAGAW-3'

5'-GATGAGTCCTGAGTAAGCW-3'

5'-GATGAGTCCTGAGTAAGTW-3'

5'-GATGAGTCCTGAGTAATAN-3'

5'-GATGAGTCCTGAGTAATCW-3'

5'-GATGAGTCCTGAGTAATGW-3'

5'-GATGAGTCCTGAGTAATTS-3'

In total, 83 Col-specific fragments were generated, of which 62 were shared by both YAC clones. Three fragments were AFLP markers polymorphic between Ws and Ler, of which two were Ws AFLP markers (a Col fragment also present in Ws and absent in Ler) and one was a Ler AFLP marker (a Col fragment also present in Ler and absent in Ws). These results are presented in Table 5 below.

Table 5 shows a number of shared and unique AFLP fragments detected in YACs 10G07 and 7E03 and the number of informative AFLP markers among these fragments in Ws and Ler genotypes.

Table 5

	AFLP fragments in YAC clones		AFLP marker	
	10G07	7E03	Ws	Ler
shared	62	62	2	1
unique	21	0	0	0

This AFLP analysis thus yielded 3 new AFLP markers (see Figure 4 and below). Their positions relative to each other and relative to the recombinant, inbred-line-derived AFLP markers were determined by analysis of the recombinants with these AFLP markers.

A second screening for AFLP markers was performed assaying all four identified YAC clones (see below) and using the enzyme combination PstI/MseI. The primers used are:

PstI-primers:

5'-GACTGCGTACATGCAGAN-3'

5'-GACTGCGTACATGCAGCW-3'

5'-GACTGCGTACATGCAGGW-3'

5'-GACTGCGTACATGCAGTN-3'

MseI-primers:

5'-GATGAGTCCTGAGTAAAN-3'

5'-GATGAGTCCTGAGTAACW-3'

5'-GATGAGTCCTGAGTAAGW-3'

5'-GATGAGTCCTGAGTAATN-3'

An "N" in the primers indicates that this part was variable (A, C, G or T) and a "W" in the primers indicates that this was A or T. A total of 144 (12 x 12) primer combinations was

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screened on all four isolated YAC clones, 12F04, 12H07, 10G07 and 7E03; the yeast strain without a YAC; and the three *Arabidopsis* ecotypes Col, Ler and Ws. In total, 219 AFLP fragments were generated, of which 144 were present in YAC clones 12F04 and 12H07 (72 were unique for clone 12F04 and 72 were shared between both YACs) and of which 75 were present in YAC clones 10G07 and 7E03 (33 were unique for clone 10G07 and 42 were shared between the 2 YACs). Three fragments derived from the first set of YAC clones were polymorphic (Ws AFLP markers). These results are presented in Table 6 below.

Table 6 lists the number of shared and unique AFLP fragments detected in YACs and the number of informative AFLP markers among these fragments in Ws and Ler genotypes.

Table 6

	number of AFLP fragments in YAC clones				AFLP markers	
	12F04	12H07	10G07	7E03	Ws	Ler
shared	72	72	0	0	1	0
unique	72	0	0	0	2	0
shared	0	0	42	42	0	0
unique	0	0	33	0	0	0

The results indicate that YAC clone 12H07 is part of the larger YAC clone 12F04, and that YAC clone 7E03 is a part of the larger YAC clone 10G07. These data indicate that the larger YAC clones, 12F04 and 10G07, do not overlap. These data do not allow the positioning of the *NIM1* gene on any of these YAC clones. The whole screening, involving 400 primer combinations producing 302 AFLP fragments in the *NIM* region, yielded 5 useful AFLP markers, of which 4 were Ws-specific and one Ler-specific. These 5 additional AFLP

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markers have been mapped by analysis of recombinant plants (see figure 4 and below) and are denominated W84.1 (a.k.a. W83.3), W84.2, W85.1, W86.1 and L86.

Table 7 lists the primer sequences used to obtain these AFLP markers. These 5 additional AFLP markers raised the total number of AFLP markers to 12 in the region from L81.1 to L88 (see figure 4 and below).

Table 7 shows primer combinations of AFLP markers derived from YAC clones.

"EcoRI-" refers to the sequence 5'-GACTGCGTACCAATTC-3',

"MseI-" refers to the sequence 5'-GATGAGTCCTGAGTAA-3' and

"PstI-" refers to the sequence 5'-GACTGCGTACATGCAG-3'.

Table 7

AFLP marker	Primer combination with selective extensions	

W84.1	PstI-AT	MseI-TT
W84.2	PstI-AA	MseI-TT
W85.1	EcoRI-CT	MseI-GTA
W86.1	EcoRI-GT	MseI-CTT
L86	EcoRI-GT	MseI-CTT

This information was used to construct a physical map of the region, as shown in Figure 5, with approximate positions of the YAC clones, relative to the genetic map. The map showed that the region containing the *NIM1* locus, between markers W83.1 and W85.1, is partly covered by 3 YAC clones: 12F04 and 10G07/7E03.

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iii. Construction of a P1/BAC contig containing the *NIM1* gene.

In the previous sections it was described how AFLP markers linked to the *NIM1* region were isolated and how YACs corresponding to these markers were identified and mapped. The results obtained while localizing the *NIM1* gene to a chromosome fragment, did not allow the definition of a specific DNA segment containing the *NIM1* gene: the flanking AFLP markers were mapped to different YACs that did not overlap. It was, therefore, not possible to determine the precise physical position of the *NIM1* gene; it could be located on either of the two YACs or in the gap between the YACs. An alternative approach was selected to close the physical gap between the flanking markers: a P1 and BAC library were employed to bridge the gap between the flanking AFLP markers.

The libraries used for gap closure were an Arabidopsis ecotype Columbia P1 library described by Liu et al (The Plant J. 7, 351-358, 1995) and an ecotype Columbia BAC library described by Choi et al (<http://genome-www.stanford.edu/Arabidopsis/ww/Vol2/choi.html>). The P1-library consists of about 10,000 clones with an average insert size of 80 kb and the BAC library consists of about 4000 clones with an average insert size of 100 kb. In theory these libraries represent about 10 nuclear genome equivalents (assuming a haploid genome size for *Arabidopsis* of 120 Mb).

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iv. Identification of P1 clones corresponding to the flanking markers.

The flanking markers Ws84.2 and Ws85.1 were used to screen pools of P1 clones using a similar strategy as previously described for screening of the YAC library (see Example 1.5i). P1 clones having the marker fragments were selected and "plasmid" DNA was isolated. The various P1 clone DNAs were fingerprinted using the ECs EcoRI/MseI and HindIII/MseI and primers without selective nucleotides. A physical map was constructed, i.e. a map giving the size and overlaps of the clones, by comparing the AFLP fingerprints. The number of AFLP fragments that are unique and the number of AFLP fragments that are common between clones indicate the extent of the overlaps. The map is displayed in Figure 6. The AFLP fingerprinting revealed that two sets of non-overlapping P1-contigs had been constructed each containing one of the flanking markers: P1-1 and P1-2 containing marker Ws84.2; P1-3 and P1-4 containing marker W85.1. Consequently, the gap between the flanking markers was not closed (Figure 6).

The positions of the P1 contigs with respect to the YAC contig was determined by AFLP fingerprinting of the YACs and P1 clones with a number of YAC-specific PCs described above. P1 clones P1-1 and P1-2 appeared to overlap completely with YAC CIC12F04, but only partially with YAC CIC12H07. Therefore, the latter P1 clones could be positioned on the YAC contig CIC12H07/12F04 (Figure 6). P1 clones P1-3 and P1-4 overlapped completely with both YACs CIC7E03 and CIC10G07 and it appeared that AFLP marker W86.1, like W85.1, was mapped to this P1 contig (Figure 6).

Next, marker L85 was used to identify corresponding P1 and BAC clones. L85 is an ecotype-Landsberg-specific marker and, therefore, colony hybridization of radioactively labelled L85 DNA to P1 and BAC filters was employed. Not a single P1 or BAC clone hybridizing to L85 was identified. This supported our earlier findings that the L85 sequence

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is lacking in the *Arabidopsis* ecotype Columbia genome and is, therefore, the most likely explanation of why no corresponding clones were identified.

v. Extending the *NIM1*-flanking P1 Contigs.

Various approaches were employed to extend from the flanking P1 contigs:

YAC AFLP fragments specific to the South end of YAC CIC12F04 (unique to CIC12F04, not present in CIC 12H07) were used to identify P1 clones by AFLP screening of pools of the library.

1. YAC AFLP fragments from YAC 10G07 and overlapping with P1-4 were used to identify P1 clones by AFLP screening of pools of the P1 library.
2. EcoRI restriction fragments from P1 clone P1-6 (resulting from the AFLP-based P1 library screening of step 1) were used as hybridization probes on filters of the BAC library.

Various P1 and BAC clones resulted from this screening and all were AFLP-fingerprinted with the ECs EcoRI/MseI and HindIII/MseI using primers without selective nucleotides. A new map was constructed as described above and is depicted in Figure 7. Table 8 shows the various AFLP PCs having AFLP fragments mapped to flanking YACs and used to screen the P1-library for corresponding P1 clones.

Table 8 represents the various AFLP PCs used to screen the P1 library. The top half of the table shows PCs specific for the North YACs and the bottom half shows the PCs specific for the South YACs. Also indicated are the YACs and P1 clones wherein the AFLP fragments were detected.

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Table 8

<u>AFLP PCs</u>	<u>CIC YACs</u>	<u>P1-clones</u>	<u>Comments</u>
PstI-AA MseI-TT	12F04 and 12H07	P1-1, P1-2	Marker Ws84.2
PstI-AT MseI-GT	12F04-specific	P1-1, P1-2	
PstI-CA MseI-AA	12F04-specific	P1-6	
PstI-AC MseI-TG	12F04-specific	P1-7	
PstI-AG MseI-TG	12F04-specific	P1-7	
PstI-CT MseI-GT	12F04-specific	P1-7	
EcoRI-CT MseI-GTA	10G07 and 7E03	P1-3, P1-4	Marker Ws85.1
EcoRI-GT MseI-CTT	10G07 and 7E03	P1-4	Marker Ws86.1
EcoRI-AA MseI-GT	10G07 and 7E03	P1-4, P1-9	
EcoRI-AT MseI-GA	10G07 and 7E03	P1-4, P1-9	
EcoRI-GG MseI-CT	10G07 and 7E03	P1-4, P1-9	

A

P1/BAC contig of about 250 kb was obtained covering the South end of YAC CIC12F04 (not extending from this YAC) and containing marker W84.2. A P1 contig of about 150 kb containing markers W85.1 and W86.1 was obtained; this contig is completely contained within YAC CIC7E03.

Construction of a P1/BAC contig covering the *NIM1* gene AFLP marker analysis on the recombinants with markers from the South end of the North P1/BAC contig (WL84.4 and WL84.5, see below and table 11) showed that the previous "walking" steps were unsuccessful in the construction of a contig containing the *NIM1* gene (see next section). Therefore, the existing North P1/BAC contig was extended South with the purpose of "walking" across the *NIM1* gene, which would enable the definition and isolation of a specific DNA segment containing the *NIM1* gene. A hybridization-based approach was followed in which P1 or BAC clones located at the South end of the North P1/BAC contig

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were used to identify clones positioned closer to *NIM1* (South bound). New clones resulting from the walking steps were mapped with respect to the existing contigs using AFLP fingerprinting with the ECs EcoRI/MseI and HindIII/MseI as described above. A total of 5 subsequent walking steps appeared to be necessary to "cross" the *NIM1* gene. Table 9 shows the clones obtained in the various walking steps.

Table 9 is an overview of the various walking steps showing the hybridization probe used to screen the P1 and BAC libraries and the selected clones hybridizing to the probes and extending in the South direction.

Table 9

	<u>Probe</u>	<u>New clones extending South</u>
Step 1:	P1-7	BAC-02
Step 2:	BAC-02	P1-16, BAC-03
Step 3:	BAC-03	P1-17, P1-18
Step 4:	P1-18	P1-21, P1-20, BAC-04
Step 5:	BAC-04	P1-22, P1-23, P1-24, BAC-06, BAC-05

A physical map of the various clones resulting from this walking effort is depicted in Figure 8. A total distance of about 600 kb was covered starting from the initial walking point marker W84.2. The South end of the contig presented in Figure 8 appeared to contain the *NIM1* gene (see next section). The contig extends more than 300 kb South from YAC CIC12F04 and appeared not to overlap with YACs CIC10G07 and CIC7E03, indicating that the *NIM1* gene is in the gap between the flanking YAC contigs and that this gap is at least 300 kb.

vi. Construction of an Integrated Genetic and Physical Map of the *NIM1* Region.

In the previous sections it was described how AFLP markers linked to the *NIM1* region were isolated, how YACs corresponding to the flanking markers were identified, and how a P1/BAC contig was constructed extending about 550 kb South from the closest North flanking AFLP marker W84.2. This section describes the generation of new AFLP markers from the P1/BAC contig, the physical mapping of these markers on this contig and the genetic mapping of these markers with the available recombinants.

1. Generation of New AFLP Markers from the P1/BAC Contig

As described in the previous section, the P1 and BAC clones of the contig extension were characterized by AFLP fingerprinting using the ECs EcoRI/MseI and HindIII/MseI. This defined quite accurately the extent of the overlaps between the various P1 and BAC clones and, in addition, generated a number of AFLP fragments specific for these clones. AFLP primers without selective nucleotides are used in fingerprinting of purified plasmid DNA of the P1 or BAC clones. Selective nucleotides will be necessary, however, to be able to use these P1 or BAC-specific AFLP fragments for detection in *Arabidopsis*. By determining the end sequences of the amplified restriction fragments, AFLP primers having the appropriate selective bases can be designed to amplify the P1- or BAC-specific AFLP fragment in *Arabidopsis*. All AFLP fragments originate from the ecotype Columbia (Col) and, therefore, it also should be determined if the Columbia AFLP markers are informative in the *NIM1* recombinants which are derived from a cross of the ecotypes Landsberg erecta (Ler) and a *nim1* mutant of the ecotype Wassilewskija (Ws-*nim*). In principle, there are 4 types of AFLP fragments, two of which are useful markers as indicated in Table 10 below:

Table 10 is an overview of the types of AFLP markers found. (+) or (-) indicates presence or absence of the AFLP fragment.

Table 10

Col	Ler	Ws-nim	marker-type
+	+	+	not informative
+	+	-	Ler marker
+	-	+	Ws marker
+	-	-	not informative

In general, fingerprinting of the P1 and BAC clones generated 30 to 40 EcoRI/MseI AFLP fragments and 60 to 80 HindIII/MseI AFLP fragments for each individual clone. The end sequences of individual fragments were determined by standard sequencing techniques. Next, specific AFLP primers sets with selective extensions of 3 nucleotides for both the EcoRI or HindIII primer and the MseI primer were tested on the following panel of DNAs:

1. P1/BAC clone from which the AFLP marker was derived
- 2a. Yeast
- 2b. YAC clone CIC12F04 (only for AFLP fragments from P1-7)
- 2c. YAC clone CIC10G07
- 3a. Col, origin of the P1 and BAC libraries
- 3b. Ler, parent 1 of the *nim* recombinants
- 3c. Ws-nim, parent 2 of the *nim* recombinants

Six clones were selected for sequence analyses of their EcoRI/MseI and HindIII/MseI AFLP fragments: BAC-01/P1-7, P1-17/P1-18, BAC-04/BAC-06. The AFLP fragments from clone P1-7 were all detected in YAC CIC12F04, indicating that this clone is completely contained within this YAC. None of the P1/BAC-specific AFLP fragments was detected in YAC clone CIC10G07, indicating that the P1/BAC contig does not bridge the gap between the two flanking YAC contigs. AFLP markers selected for analysis of the *nim* recombinants are depicted in Table 11.

Table 11 is an overview of the selected AFLP markers from the AFLP PCs specific for the various P1 and BAC clones. A "WL" marker is a marker originating from the same PC and displaying two AFLP markers, a Ws and a Ler marker, which appeared to be completely linked in repulsion phase upon analysis of the *NIM* recombinants.

Table 11

<u>Origin</u>	<u>Marker name</u>	<u>AFLP primers combination</u>
P1-7	WL84.4	EcoRI-AGC MseI-ACT
P1-7	WL84.5	HindIII-CTC MseI-TTC
P1-17/P1-18	Ler84.6a	HindIII-CGT MseI-ATT
P1-17/P1-18	Ler84.6b	HindIII-ATT MseI-CAT
P1-18	Ler84.6c	HindIII-TCT MseI-TAT
P1-18	Ler84.7	EcoRI-AAA MseI-AGA
BAC-04/06	Ler84.8	EcoRI-TTC MseI-AGT
BAC-06	Ler84.9a	EcoRI-AAA MseI-TGT
BAC-06	Ler84.9b	EcoRI-ATC MseI-TCC
BAC-06	Ler84.9c	EcoRI-ATG MseI-GTA

2. Physical Mapping of the New AFLP Markers.

The AFLP markers described above were physically mapped by detecting their presence in the various P1 and BAC clones. The results are presented in Figures 9-11.

3. Genetic Mapping of the New AFLP Markers.

The AFLP markers were all analyzed on a selected set of the recombinants. The results obtained are summarized in Tables 12a, 12b and 12c.

Table 12a
NIM RECOMBINANTS NORTH OF WL84.4&5

Nr.	Plant	PR- 1 on/ off	Ler 84	Ws 84. 2	WL 84. 4& 5	Ler 84. 6a	Ler 84. 6b	Ler 84. 6c	Ler 84. 7	nim	Ler 85	Ler 84. 8	Ler 84. 9b	Ler 84. 9a	Ler 84. 9c	Ws 85. 1	Ws 86. 1	Ler 86
N1	A-74 nim	off	(H)	H	W			W	W	W	W	W			W	W	W	W
N2	A-113		H	H	W			W	W	W	W	W			W	W	W	W
N3	B-023 Rnim	off	H	H	W			W	W	W	W	W			W	W	W	W
N4	B-297 notnim	on	W	W	H	(H)	H	H	H	H	H	H	H	H	H	H	H	H
N5	B-292 nim	off	H	H	W			W	W	W	W	W			W	W	W	W
N6	D-269	off	H	H	W			W	W	W	W	W			W	W	W	W

	W	W	W	W	W	W	W	W
	W	W	W	W	W	W	W	W
	W	W	W	(H)	W	W	W	W
	W	W	W		W		W	W
						W		
						W		
	W	W	W		W	W	W	W
	W	W	W		W	W	W	W
	W	W	W	W	W	(W)	W	W
	W	W	W	W	W		W	W
	W	W	W	W	W		W	W
						W		
						W		
	W	W	W		W	W	W	W
	H	H	H	H	H	H	H	H
	H	H	H	H	(H)	H	H	H
	off	off	off	off	off	off	off	off
Rnim	D-306 nim	E-086 nim	F-049 Rnim	G-002 nim	G-009 Rnim	G-064 (nim)	G-072 nim	H-037 nim
N7								
N8								
N9								
N10								
N11								
N12								
N13								
N14								

N15	H-047 Rnim	off	H	H	W					W	W
N16	H-097 (nim)	off	H	H	W	W	W	W	W	W	W

Table 12b
NIM RECOMBINANTS SOUTH OF Ler84.9c

Nr.	Plant	PR- 1 on/ off	Ler 84	Ws 84. 2	WL 84. 4& 5	Ler 84. 6a	Ler 84. 6b	Ler 84. 6c	Ler 84. 7	nim	Ler 85	Ler 84. 8	Ler 84. 9b	Ler 84. 9a	Ler 84. 9c	Ws 85. 1	Ws 86. 1	Ler 86
S6	A-116 Rnim	off	W	W	W			W	W	W	W	W	W	W	W	H	H	H
S7	B-111 nim	off	W	W	W			W	W	W	W	W	W	W	W	H	H	H
S8	B-165 notnim	inter m	H	H	H	H	H	H	H	H	H	H	H	H	H	W	W	W
S9	B-182 notnim	inter m.	H	H	H	H	H	H	H	H	H	H	H	H	H	W	W	W
S10	B-190 segWT	off	W	W	W			W	W	(W)	W	W	W	W	W	H	H	H

[illegible]

Table 12c

NIM RECOMBINANTS BETWEEN WL84.4&5 AND Ler84.9c

Nr.	Plant	PR-	Ler	Ws	WL	Ler	Ler	Ler	Ler	Ler	nim	Ler	Ler	Ler	Ler	Ler	Ws	Ws	Ler
		1	84	84.	84.	84.	84.	84.	84.	84.	85	84.	84.	84.	84.	84.	85.	86.	86
		on/		2	4&	6a	6b	6c	7								1		
		off			5														
N17	B-304 nim		H	H	H	W	W	W	W	W	W	W	W	W	W	W	W	W	W
N18	C-111 nim	off	H	H	H	W	W	W	W	W	W	W	W	W	W	W	W	W	W
N19	E-093 nim	off	H	H	H	W	W	W	W	W	W	W	W	W	W	W	W	W	W
N20	E-110 Rnim	off	H	H	H	W	W	W	W	W	W	W	W	W	W	W	W	W	W
N21	G-014 nim	off	H	H	H	W	W	W	W	W	W	W	W	W	W	W	W	W	W
N22	A-019	int	W	W	W	W	H	H	H	H	H	H	H	H	H	H	H	H	H

		erm	notnim	
	H	H	H	H
	H	H	H	H
	H	H	H	H
	H	H	H	H
	H	H	H	H
	H	H	H	H
	H	H	H	H
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	H	H	H	H
	H	H	H	H
	H	H	H	H
	H	H	H	H
	H	H	H	H
	H	H	H	H
	H	H	H	H
	H	H		

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The AFLP markers Ler84.8, Ler84.9a, Ler84.9b and Ler84.9c appeared to map at the South side of *NIM1*. Recombinants were found that were phenotypically *nim1* (homozygous, genotype *Ws-nim1/Ws-nim1*) and heterozygous for these AFLP markers (the Ler-specific AFLP marker was detected, genotype is *Ws-nim1/Ler*). AFLP marker Ler84.8 appeared to be closest to *NIM1*: only a single recombinant (C-105) was scored as heterozygous *Ws-nim1/Ler* and homozygous *Ws-nim1/Ws-nim1*. AFLP markers Ler84.7 and Ler84.6c appeared to completely cosegregate with *NIM1*: all recombinants had an identical *NIM1* and AFLP marker genotype. North of *NIM1*, marker L84.6b appeared to be closest to *NIM1*: three *nim1* phenotype recombinant plants, C-074, D-169 and E-103 (Table 12c), were found to be heterozygous *Ws-nim1/Ler* at this marker. With the aid of the cosmid contig generated from P1-18, BAC-04 and BAC-06, AFLP markers Ler84.6b and Ler84.8 were mapped in P1-18 and BAC-04, respectively, and found to have a physical distance of approximately 110 kb. This defines *nim1* to be located on a DNA segment estimated to be 110 kb in length. From this analysis it has been determined that the *NIM1* gene is contained in clone BAC-04 or P1-18. Clones BAC-04 and P1-18 have been deposited with ATCC and given deposit numbers ATCC 97543 and ATCC 97606, respectively.

vii. Genetic and Physical Fine Mapping of the *NIM1* Gene.

The previous section described how a DNA segment containing the *NIM* gene was delineated by physical mapping of the flanking AFLP markers (Ler84.6b and Ler84.8) on the P1/BAC contig. The flanking markers appeared to map on two overlapping clones, P1-18 and BAC-04. This section describes how additional BAC-04-specific and P1-18-specific AFLP markers were generated to increase the resolution of the genetic and physical map in the region containing the *NIM1* gene.

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viii. Generation of Additional AFLP Markers from the Cosmid Array.

Four ECs were selected to generate additional AFLP markers for fine mapping of *NIM1*: PstI/MseI, XbaI/MseI, BstYI/MseI and TaqI/MseI. PstI/MseI and XbaI/MseI AFLP fragments were generated on clone P1-18 and BAC-04 and the selective sequences necessary for detection in *Arabidopsis* were determined. Similarly, the AFLP fragments and selective sequences were determined for BstYI/MseI and TaqI/MseI; however, in this case the procedure was performed using cosmid DNAs: A11, C7, E1 and E8 for BstYI/MseI (complete *NIM1* region) and D7, E8 and E6 for TaqI/MseI (South side of *NIM1* region). Informative AFLP markers selected for further genetic and physical mapping are shown in Table 13. Additional adapters used in this work are shown in Table 14.

Table 13 shows the AFLP markers used for genetic and physical fine mapping of *NIM1*. "BstYI(T)" indicates that the restriction site and corresponding primer was either AGATCT or GGATCT.

Table 13

Marker	EC/PC	
Ler84.Y1	BstYI(T)-GCT	MseI-AAC
Ws84.Y2	BstYI(T)-TCT	MseI-GCA
Ler84.Y3	BstYI(T)-AAG	MseI-TAT
Ler84.Y4	BstYI(T)-GTT	MseI-AGA
Ws84.T1	TaqI-TAC	MseI-GGA
Ler84.T2	TaqI-TTG	MseI-GGA

Table 14 shows same Additional adapters used for identifying new AFLP markers.

Table 14

BstYI: 5'-CTCGTAGACTGCGTACC-3'
3'-CATCTGACGCATGGCTAG-5'

TaqI: 5'-CTCGTAGACTGCGTACC-3'
3'-CATCTGACGCATGGGC-5'

ix. Physical mapping of new AFLP markers to the cosmid contig.

The markers indicated above were physically mapped on the cosmid array by determining their presence in the various cosmid clones (Figure 11).

1. Genetic Mapping of New AFLP Markers.

The new AFLP markers were genetically mapped by AFLP analysis of the closest North and South recombinants. The closest North (recombinant D169) and South (recombinant C105) recombination points were mapped (see Table 15). The AFLP analysis showed that recombinant D169 had a recombination South of marker L84.Y1, but North of marker W84.Y2. The recombination point in recombinant C105 mapped between markers L84.T2 and L84.8. Using the available set of recombinants this allowed further delineation of the chromosomal interval containing *NIM1*; the distance between the flanking recombination points appeared to be 60-90 kb (Figure 12).

Table 15 NIM RECOMBINANTS BETWEEN WL84.4&5 AND Ler84.9c

[illegible]

56/2

H	W	W	W	H	H	H	H
H	W	W	W	H	H	H	H
H	W	W	W	H	H	H	H
H	W	W	W	H	H	H	H
H	W	W	W	H	H	H	W
H	W	W	W	H	H	W	W
H	W	W	W	H	W	W	W
H	W	W	W	W	W	W	W
H	W	W	W	W	W	W	W
H	W	W	W	W	W	W	W
H	W	W	W	W	W	W	W
H	W	W	W	W	W	W	W
H	W	W	W	W	W	W	W
H	W	W	W	W	W	W	W
H	W	W	W	W	W	W	W
H	W	W	W	W	W	W	W
H	W	W	H	W	W	W	W
H	H	H	H	W	W	W	W
W	H	H	H	W	W	W	W
W	H	H	H	W	W	W	W
W	H	H	H	W	W	W	W
W	H	H	H	W	W	W	W
int	off	off	off	off		off	off
A-019 notni m	C-074 Rnim	E-103 nim	D-169 nim	C-105 nim	H-039 nim	B-052 nim	B-142 nim
N22	N23	N24	N25	S1	S2	S3	S4

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2. Construction of a Cosmid Contig.

For complementation of the *nim1* plant phenotype, transformation of *nim1* plants is required with a wildtype *NIM1* gene. This can be accomplished by transforming these plants with a cosmid containing the gene. For this purpose, a cosmid contig of the *NIM1* region is constructed. Since *Arabidopsis* is transformed using *Agrobacterium*, the cosmid vector used is a binary vector.

DNA was isolated from BAC-04, BAC-06 and P1-18, and used to make a partial digest using restriction enzyme *Sau3AI*. The 20-25 kb fragments were isolated using a sucrose gradient, pooled, and filled in with dATP and dGTP. The binary vector (04541) was cleaved with *XhoI* and filled in with dCTP and dTTP. The fragments were next ligated into the vector. The ligation mix was packaged and transduced into *E.coli*.

This cosmid library was screened with the BAC-04, BAC-06 and P1-18 clones and positive clones isolated. These cosmids were next AFLP fingerprinted and arranged into a contig of overlapping clones spanning the *NIM1* region. The insert sizes of the cosmids were determined, and limited restriction mapping was performed. The results are shown in Figure 10.

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Example 2

Identification of a Clone Containing the *NIM1* gene.

1. Complementation Via Stable Transformation

Cosmids that are generated from clones that span the *NIM1* region (described above) are moved into *Agrobacterium* by triparental mating. These cosmids are then used to transform *nim1 Arabidopsis* by vacuum infiltration (Mindrinos et al., 1994, Cell 78, 1089-1099) or by standard root transformation. Seed from these plants is harvested and allowed to germinate on agar plates with kanamycin (or another appropriate antibiotic) as selection agent. Only plantlets that are transformed with cosmid DNA can detoxify the selection agent and survive. Seedlings that survive the selection are transferred to soil and tested for the *nim* phenotype or their progeny are tested for the *nim* phenotype. Transformed plants that no longer have the *nim* phenotype identify cosmid(s) that contain a functional *NIM1* gene.

2. Complementation in a Transient Expression System.

The ability of DNA clones to complement the *nim1* mutation is tested in 2 transient expression systems.

In the first system, *nim1 Arabidopsis* plants containing a PR1-luciferase (PR1-lux) transgene are used as bombardment recipient material. These plants are generated by transforming Columbia ecotype plants with a PR1-lux construct by vacuum infiltration, followed by kanamycin selection of the harvested seed, as described above. Transformed plants that express luciferase activity after induction with INA are selfed and homozygous plants are generated. These are crossed to *nim1* plants. In the transient assay, progeny plants from this cross that are homozygous for *nim1* and for PR1-lux are used for identification of DNA clones that can complement the *nim1* phenotype. To this end, the plants are first treated with INA, as described in example 1.1 above. Two days later these

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plants are harvested, surface sterilized and plated on GM agar medium. The leaf tissue is then bombarded with cosmid, P1 or BAC clones (or subclones) from the *NIM1* region and after one day, the luciferase activity of the leaves is measured. Clones that induce luciferase activity contain the *NIM1* gene.

In a second system, *nim1* plants are treated with INA (as described in example 1.1 above) and 2 days later bombarded with cloned DNA (cosmid, P1, BAC and/or YAC clones or subclones) from the *NIM1* locus region and a reporter plasmid. The reporter plasmid contains the luciferase gene, driven by the *Arabidopsis* PR1 promoter (PR1-lux). In *nim1* plants, INA does not activate the PR1 promoter (as described in example 1.2 above) and thus can not induce luciferase activity from the reporter plasmid. However, when a cotransformed DNA clone contains the complementing *NIM1* gene, INA does induce the PR1 promoter, as evidenced by an induction of luciferase activity. One day after the cobombardment, the luciferase activity of the whole plant is measured. DNA clones (cosmids, P1 or BAC clones or subclones) that induce luciferase activity that is significantly above background levels contain the *NIM1* gene.

3. Changes in transcripts in *nim1* phenotype lines.

Since *nim1* phenotype plants have mutations in the *NIM1* gene, it is conceivable that in some lines the gene is altered in such a manner that there is no mRNA transcribed, or an aberrant mRNA (size) is produced. To test for this, RNA blot analysis is performed on the *nim1* lines.

RNA is isolated from Ws and Ler plants of these lines, (after water or INA or BTH treatment) and used to prepare northern blots. These blots are hybridized with DNA fragments isolated from clones of the DNA contig of the *NIM1* locus. DNA fragments that identify *nim1* lines with aberrant RNA expression (aberrant in size or concentration), likely

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identify (part of) the *NIM1* gene. The DNA fragment and surrounding DNA is sequenced and used to isolate a cDNA (by library screen or by reverse transcription-PCR), which is also sequenced. The clone from which the fragment was isolated or the isolated cDNA is used to show complementation of the *nim1* phenotype in stable and transient expression systems.

Example 3

Determination of the DNA sequence of the *NIM1* gene.

1. Genomic sequencing.

Genomic clones that may contain the *NIM1* gene are sequenced using methods known in the art. These include BAC-04, P1-18 and the cosmids from the *NIM1* region. For instance, the cosmids are digested with restriction enzymes and fragments that are derived from the insert are cloned into a general purpose vector, such as pUC18 or Bluescript. The larger P1 and BAC clones are randomly sheared and fragments cloned into a general purpose vector. The fragments in these vectors are sequenced by conventional methods (e.g. by "primer walking" or generation of deletions of inserts). The obtained sequences are assembled into a contiguous sequence.

The sequence of the insert of a complementing clone contains the *NIM1* gene. The approximate start and end of the *NIM1* gene are deduced based on the DNA sequence, sequence motifs such as TATA boxes, the open reading frames present in the sequence, codon usage, the cosmid complementation data, the relative location of the AFLP markers and additional relevant data that is gathered (see Example 4, below).

2. cDNA sequencing.

The cosmid(s) or larger clones that contain the *NIM1* gene (as described in Example 2 above), are used to isolate cDNAs. This is accomplished by using the clones (or DNA fragments) as probes in a screen of a cDNA library of wildtype *Arabidopsis* plants. The cDNAs that are isolated are sequenced as described for cosmid sequencing and used in complementation tests. To this end, full length cDNAs are cloned into a suitable plant expression vector, behind a constitutive promoter. These constructs are used in the transient assays as described above. Alternatively, the cDNAs are cloned into a binary expression vector, allowing for expression in plant tissues and for *Agrobacterium*-mediated plant transformation, as described in Example 2 above. A cDNA that contains the *NIM1* gene (as determined by complementation, isolation with a closely linked AFLP marker, isolation with a cosmid fragment, or by other deduction) is sequenced.

The genes from Ws-O and *nim1* plants are isolated and sequenced. The genes are obtained from a cosmid or cDNA library, using a fragment of the isolated *NIM1* gene as a probe. Alternatively, the genes or cDNAs are isolated by PCR, using *NIM1*-gene-specific primers and genomic DNA or cDNA as template. Likewise, the *nim1* alleles from other *nim1* lines (see Example 1.1 above) are isolated and sequenced in a similar manner.

Example 4

Description of the *NIM1* gene and deduced protein sequence

The DNA sequence of the *NIM1* gene or cDNA is determined as described in Example 3 above. This sequence is analyzed with the use of DNA analysis programs, such as can be found in the Genetics Computer Group (GCG) package, in the Sequencer or Staden packages, or any similar DNA analysis program package.

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Specifically, the start and end of the gene are determined, based on open reading frame analysis, the presence of stop and potential start codons, the presence of potential promoter motifs (such as the TATA box), the presence of polyadenylation signals and the like. Also, the predicted amino acid is deduced from the open reading frame. Both the DNA and protein sequence are used to search databases for sequences with homologies, such as transcription factors, enzymes or motifs of such genes or proteins.

Example 5

Isolation of *NIM1* homologs

The *Arabidopsis NIM1* gene may be used as a probe in the low stringency hybridization screening of a genomic or cDNA library in order to isolate *NIM1* homologs from other plant species. Alternatively, this is accomplished by PCR amplification, using primers designed based on the *Arabidopsis NIM1* gene sequence and using genomic DNA or cDNA as template. The *NIM1* gene may be isolated from corn, wheat, rice, barley, rape seed, sugarbeet, potato, tomato, bean, cucumber, grape, tobacco and other crops of interest and sequenced. With a set of sequences from *NIM1* gene homologs in hand, new primers can be designed from conserved portions of the gene, in order to isolate *NIM1* homologs from more distantly related plant species by PCR amplification.

Example 6

Complementation of the *nim1-1* gene with genomic fragments.

1. Construction of a cosmid contig.

A cosmid contig of the *NIM1* region was constructed using CsCl-purified DNA from BAC04, BAC06 and P1-18. The DNAs of the three clones were mixed in equimolar quantities and were partially digested with the restriction enzyme Sau3A. The 20-25 kb

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fragments were isolated using a sucrose gradient, pooled and filled in with dATP and dGTP. Plasmid pCLD04541 was used as T-DNA cosmid vector. This plasmid contains a broad host range pRK290-based replicon, a tetracycline resistance gene for bacterial selection and the *nptII* gene for plant selection. The vector was cleaved with *Xho*I and filled in with dCTP and dTTP. The prepared fragments were then ligated into the vector. The ligation mix was packaged and transduced into *E. coli* strain XL1-blue MR (Stratagene). Resulting transformants were screened by hybridization with the BAC04, BAC06 and P1-18 clones and positive clones isolated. Cosmid DNA was isolated from these clones and template DNA was prepared using the ECs *Eco*RI/*Mse*I and *Hind*III/*Mse*I. The resulting AFLP fingerprint patterns were analyzed to determine the order of the cosmid clones. A set of 15 semi-overlapping cosmids was selected spanning the *nim* region (Figure 13). The cosmid DNAs were also restricted with *Eco*RI, *Pst*I, *Bss*HII and *Sgr*AI. This allowed for the estimation of the cosmid insert sizes and the verification of the overlaps between the various cosmids as determined by AFLP fingerprinting.

2. Identification of a clone containing the *NIM* gene.

Cosmids generated from clones spanning the *NIM1* region were moved into *Agrobacterium tumefaciens* AGL-1 through conjugative transfer in a tri-parental mating with helper strain HB101 (pRK2013). These cosmids were then used to transform a kanamycin-sensitive *nim1-1* *Arabidopsis* line using vacuum infiltration (Mindrinos et al., 1994, Cell 78, 1089-1099). Seed from the infiltrated plants was harvested and allowed to germinate on GM agar plates containing 50 mg/ml kanamycin as a selection agent. Only plantlets that are transformed with cosmid DNA can detoxify the selection agent and survive. Seedlings that survive the selection were transferred to soil approximately two weeks after plating and

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tested for the *nim1* phenotype as described below. Transformed plants that no longer have the *nim1* phenotype identify cosmid(s) that contain a functional *NIM1* gene.

3. Testing for the *nim1* phenotype of transformants.

Plants transferred to soil were grown in a phytotron for approximately one week after transfer. 300µm INA was applied as a fine mist to completely cover the plants using a chromister. After two days, leaves were harvested for RNA extraction and PR-1 expression analysis. The plants were then sprayed with *Peronospora parasitica* (isolate EmWa) and grown under high humidity conditions in a growing chamber with 19°C day/17° night temperatures and 8h light/16h dark cycles. Eight to ten days following fungal infection, plants were evaluated and scored positive or negative for fungal growth. Ws and *nim1* plants were treated in the same way to serve as controls for each experiment.

Total RNA was extracted from the collected tissue using a LiCl/phenol extraction buffer (Verwoerd, et al. NAR 17:2362). RNA samples were run on a formaldehyde agarose gel and blotted to GeneScreen Plus (DuPont) membranes. Blots were hybridized with a ³²P-labeled PR-1 cDNA probe. The resulting blots were exposed to film to determine which transformants were able to induce PR-1 expression after INA treatment. The results are summarized in Table 16.

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Table 16 shows complementation of the *nim1* phenotype by cosmid clones.

Table 16

Clone Name	# of transformants	# of plants with INA induced PR-1/total # of plants tested (%)
A8	3	0/3 (0%)
A11	8	4/18 (22%)
C2	10	1/10 (10%)
C7	33	1/32 (3%)
D2	81	4/49 (8%)
D5	6	5/6 (83%)
E1	10	10/10 (100%)
D7	129	36/36 (100%)
E8	9	0/9 (0%)
F12	6	0/6 (0%)
E6	1	0/1 (0%)
E7	34	0/4 (0%)
WS-control (wild-type)	NA	28/28 (100%)
<i>nim1</i> phenotype control	NA	0/34 (0%)

NA-not applicable

Example 7

Sequencing of the 9.9 Kb *NIM1* gene region.

BAC04 DNA (25 ug, obtained from KeyGene) was the source of DNA used for sequence analysis. This BAC was shown to be the clone completely encompassing the region that complemented the *nim1* mutants. DNA was randomly sheared using an

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approach from Cold Spring Harbor. Briefly, BAC DNA was sheared in a nebulizer to an average molecular weight of about 2 kb. Ends of the sheared fragments were repaired using a two-step protocol with dNTPS, T4 DNA polymerase and Klenow fragment (Boehringer). The end-repaired DNA was run on a 1% low-melt agarose gel and the region between 1.3 kb and 2.0 kb was cut from the gel. DNA was isolated from the gel fragment by a freeze-thaw approach. DNA was then mixed with EcoRV-digested pBRKanF4 and was ligated overnight at 4°C. pBRKanF4 is a derivative of pBRKanF1, which was obtained from Kolavi Bhat at Vanderbilt University (Bhat, K.S., Gene 134(1), 83-87 (1993)). *E. coli* strain DH5a was transformed with the ligation mix, and the transformation mix was plated onto plates containing kanamycin and X-gal. 1600 white or light blue KanR colonies were selected for plasmid isolation. Individual colonies were picked into 96-well deep well plates (Polyfiltronics, #U508) containing 1.5 ml of TB + Kan (50 ug/ml). Plates were covered and were placed on a rotating platform shaker at 37°C for 16 hrs. Plasmid DNA was isolated using the Wizard Plus 9600 Miniprep system (Promega, #A7000) according to manufacturer's recommendations.

Plasmids were sequenced using Dye Terminator chemistry (Applied BioSystems PRISM Dye Terminator Cycle Sequencing Ready Reaction Kit, P/N 402078) and primers designed to sequence both strands of the plasmids. Data was collected on ABI 377 DNA sequencers. Approximately 75% of these reactions yielded useful sequence information. Sequences were edited and were assembled into contigs using Sequencer 3.0 (Gene Codes Corporation), Staden gap4 (Roger Staden, e-mail address rs@mrc-lmb.cam.ac.uk), and PHRED (Phil Green, e-mail address phg@u.washington.edu). The largest contig (approximately 76 kb) covered the complementing region to an average depth of 7 independent calls/base.

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A region of approximately 9.9 kb defined by the overlap of cosmids E1 and D7 was identified by complementation analysis to contain the *nim1* region. Primers which flanked the insertion site of the vector and specific to the cosmid backbone were designed using Oligo 5.0 Primer Analysis Software (National Biosciences, Inc.). DNA was isolated from cosmids D7 and E1 using a modification of the ammonium acetate method (Traynor, P.L., 1990. BioTechniques 9(6): 676.) This DNA was directly sequenced using Dye Terminator chemistry above. The sequence obtained allowed determination of the endpoints of the complementing region.

A truncated version of the BamHI-EcoRV fragment was also constructed, resulting in a construct which contains none of the "Gene 3" region (Fig. 13). The following approach was necessary due the presence of HindIII sites in the Bam-Spe region of the DNA. The BamHI-EcoRV construct was completely digested with SpeI, then was split into two separate reactions for double digestion. One aliquot was digested with BamHI, the other HindIII. A BamHI-SpeI fragment of 2816 bp and a HindIII-SpeI fragment of 1588 bp were isolated from agarose gels (QiaQuick Gel extraction kit) and were ligated to BamHI-HindIII-digested pSGCG01. DH5a was transformed with the ligation mix. Resulting colonies were screened for the correct insert by digestion with HindIII following preparation of DNA using Wizard Magic MiniPreps (Promega). A clone containing the correct construct was electroporated into *Agrobacterium* strain GV3101 for transformation of *Arabidopsis* plants.

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Example 8

Identification of the *NIM1* gene region by allele sequencing.

Table 17. Genetic segregation of non-inducible immunity mutants.

Mutant	Generation	Phenotype			
		Female	Male	Wildtype	<i>nim1</i>
<i>nim1-1^a</i>	F1	wildtype ^b	<i>nim1-1</i>	24	0
	F2			98	32
<i>nim1-2</i>	F1	<i>nim1-2</i>	Wildtype	3	0
<i>nim1-3</i>	F1	<i>nim1-3</i>	Wildtype	3	0
<i>nim1-4</i>	F1	<i>nim1-4</i>	Wildtype	3	0
<i>nim1-5</i>	F1	<i>nim1-5</i>	Wildtype	0	35
<i>nim1-6</i>	F1	<i>nim1-6</i>	Wildtype	3	0
<i>nim1-2</i>	F1	<i>nim1-2</i>	<i>nim1-1</i>	0	15
<i>nim1-3</i>	F1	<i>nim1-3</i>	<i>nim1-1</i>	0	10
<i>nim1-4</i>	F1	<i>nim1-4</i>	<i>nim1-1</i>	0	15

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<i>nim1-5</i>	F1	<i>nim1-5</i>	<i>nim1-1</i>	0	14
	F2			9	85
<i>nim1-6</i>	F1	<i>nim1-6</i>	<i>nim1-1</i>	0	12

^a Data from Delaney et al. (1995) PNAS 92,6602-6606.

^b Wild type denotes the wildtype Ws-0 strain.

1. Genetic Analyses

To determine dominance of the various mutants that displayed the *nim1* phenotype, pollen from wildtype plants was transferred to the stigmata of *nim1-1*, -2, -3, -4, -5, -6. If the mutation is dominant, then the *nim1* phenotype will be observed in the resulting F1 plants. If the mutation is recessive, then the resulting F1 plants will exhibit a wildtype phenotype.

The data presented in Table 17 show that when *nim1-1*, -2, -3, -4 and -6 are crossed with the wildtype, the resulting F1 exhibit the wildtype phenotype. Thus, these mutations are recessive. In contrast, the *nim1-5* X wildtype F1 progeny all exhibit the *nim1* phenotype, indicating that this is a dominant mutation. Following INA treatment, no *P. parasitica* sporulation was observed on wildtype plants, while the F1 plants supported growth and some sporulation of *P. parasitica*. However, the *nim1* phenotype in these F1 plants was less severe than observed when *nim1-5* was homozygous.

To determine allelism, pollen from the kanamycin-resistant *nim1-1* mutant plants was transferred to the stigmata of *nim1-2*, -3, -4, -5, -6. Seeds resulting from the cross were plated onto Murashige-Skoog B5 plates containing kanamycin at 25 ug/ml to verify the hybrid origin of the seed. Kanamycin resistant (F1) plants were transferred to soil and assayed for the *nim1* phenotype. Because the F1 progeny of the cross of the *nim1-5*

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mutant with the *Ws* wildtype displayed a *nim1* phenotype, analysis of *nim1-5 X nim1-1* F2 was also carried out.

As shown in Table 17, all of the resulting F1 plants exhibited the *nim1-1* phenotype. Thus, the mutation in the *nim1-2*, *-3*, *-4*, *-5*, *-6* was not complemented by the *nim1-1*; these plants all fall within the same complementation group and are therefore allelic. Analysis of F2 progeny from the *nim1-5 X nim1-1* cross also displayed the *nim1* phenotype, confirming that *nim1-5* is a *nim1* allele.

2. Sequence Analysis and Subcloning of the NIM1 Region

The 9.9 kb region containing the *NIM1* region was analyzed for the presence of open reading frames in all six frames using Sequencher 3.0 and the GCG package. Four regions containing large ORF's were identified as possible genes (Gene regions 1-4). These four regions were PCR amplified from DNA of the wild-type parent and six different *nim1* allelic variants. Primers for these amplifications were selected using Oligo 5.0 (National Biosciences, Inc.) and were synthesized by Integrated DNA Technologies, Inc. PCR products were separated on 1.0% agarose gels and were purified using the QIAquick Gel Extraction Kit. The purified genomic PCR products were directly sequenced using the primers used for the initial amplification and with additional primers designed to sequence across any regions not covered by the initial primers. Average coverage for these gene regions was approximately 3.5 reads/base.

Sequences were edited and were assembled using Sequencher 3.0. Base changes specific to various *nim1* alleles were identified only in the region designated Gene Region 2.

The positions listed in Table 18 relate to Figure 14 and relate to the top strand of the 9.9 kb region featured in Figure 13. The open reading frames from the gene regions described in Figure 13 as 1, 2, 3 and 4 were sequenced and the changes in the different

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nim1 alleles are shown in the Table. The changes that are described are on the top strand, 5' to 3', as it would relate to Figure 13.

It is apparent that the *NIM1* gene was cloned and that it lies within Gene Region 2, since there are amino acid changes or alterations of sequence within the open reading frame of Gene Region 2 in all 6 *nim1* alleles. At the same time, at least one of the *nim1* alleles shows no changes in the open reading frames within Gene Regions 1, 3 and 4. Therefore, the only gene within the 9.9 kb region that could be *NIM1* is the Gene Region 2, the *NIM1* gene.

The Ws section of Table 18 indicates the changes in the Ws ecotype of *Arabidopsis* relative to the Columbia ecotype of *Arabidopsis*. Figures 13, 14, 15 and all others wherein sequence is shown relate to the Columbia ecotype of *Arabidopsis*, which contains the wild type gene in the experiments that were conducted. The changes are listed as amino acid changes within the gene 2 or *NIM1* region and are listed as changes in base pairs in the other regions.

Figure 13 shows 4 different panels that describe the cloning of the *NIM1* gene and describe the entire 9.9 kb region. Figure 14 is the sequence of the entire 9.9 kb region in the same orientation as described in Figure 13. Figure 15 is the sequence of the specific *NIM1* gene region which is gene region 2 indicated in Figure 13; the sequence of Figure 15 contains the *NIM1* gene. Figure 15 shows the amino acid sequence in single letter code and shows the full length cDNA and RACE product that was obtained in capital letters in the DNA sequence. Some of the allele mutations that were found are shown above the DNA sequence and the particular *nim1* allele that had that change is indicated.

Sequence analysis of the region and sequencing of various *nim1* alleles (see below) allowed identification of a cosmid region that contains the *nim1* gene. This region is delineated by a BamH1-EcoRV restriction fragment of ~5.3 kb. Cosmid DNA from D7 and

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plasmid DNA from pBlueScriptII(pBSII) were digested with Bam HI and with EcoRV (NEB). The 5.3 kb fragment from D7 was isolated from agarose gels and was purified using the QIAquick gel extraction kit (# 28796, Qiagen). The fragment was ligated overnight to the Bam-EcoRV-digested pBSII and the ligation mixture was transformed into *E. coli* strain DH5a. Colonies containing the insert were selected, DNA was isolated, and confirmation was made by digestion with HindIII. The Bam- EcoRV fragment was then engineered into a binary vector (pSGCG01) for transformation into *Arabidopsis*.

3. Northern analysis of the four gene regions.

Identical Northern blots were made from RNA samples isolated from water-, SA-, BTH- and INA-treated Ws and *nim1* lines as previously described (Delaney et al, 1995, PNAS 92, 6602-6606). These blots were hybridized with PCR products generated from the four gene regions identified in the 9.9 kb *NIM1* gene region. Only the gene region containing the *NIM1* gene (Gene Region 2) had detectable hybridization with the RNA samples, indicating that only the *NIM1* region contains a detectable transcribed gene (Figure 16 and Table 18).

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Table 18 shows *nim1* allele sequence variation.

Table 18

Allele/ ecotype	Gene Region			
	1 (bases 590- 1090)	2 (<i>NIM1</i>) (bases 1380-4100)	3 (bases 5870 - 6840)	4 (bases 8140- 9210)
<i>nim1-1</i>	no changes	t inserted at 2981: change of 7AA and premature termination of protein.	no changes	no changes
<i>nim1-2</i>	no changes	g to a at 2799: His to Tyr	no changes	no changes
<i>nim1-3</i>	no changes	deletion of t at 3261: change of 10AA and premature termination of protein.	no changes	no changes
<i>nim1-4</i>	no changes	c to t at 2402: Arg to lys	no changes	no changes
<i>nim1-5</i>	no changes	c to t at 2402: Arg to lys	no changes	no changes
<i>nim1-6</i>	g to a at 734: asp to lys	g to a at 2670: Gln to Stop	no changes	no changes
WS (compared to Columbia)	no changes	a to g at 1607: Ile to Leu a to c at 2344: intron t to g at 2480: Gln to Pro g to c at 2894: Ser to Trp ggc deleted at 3449: lose Ala c to t at 3490: Ala to Thr c to t at 3498: Ser to Asn a to t at 3873: non-coding g to a at 3992: non-coding	t to a at 5746 a to t at 5751 t to a at 5754 c to t at 6728 a to t at 6815 t to c at 6816	t to g at 8705 g to t at 8729 g to t at 8739 g to t at 8784 c to a at 8789 c to t at 8812 a to g at 8829 t to g at 8856 a to c at 9004

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		g to a at 4026: non-coding g to a at 4061: non-coding		a to t at 9011 a to g at 8461
RNA detected	No	Yes	No	No

Positions listed in the table relate to Figure 14 containing the 9.9Kb sequence. All alleles *nim1-1* to *nim1-6* are WS strain. Columbia-0 represents the wild type

We also demonstrated that the gene region 2 (Fig. 13) contains the functional *NIM1* gene by doing additional complementation experiments. A BamHI/HindIII genomic DNA fragment containing gene region 2 was isolated from cosmid D7 and was cloned into the binary vector pSGCG01 containing the gene for kanamycin resistance (Fig. 13; Steve Goff, personal communication). The resulting plasmid was transformed into the *Agrobacterium* strain GV3101 and positive colonies were selected on kanamycin. PCR was used to verify that the selected colony contained the plasmid. Kanamycin-sensitive *nim1-1* plants were infiltrated with this bacteria as previously described. The resulting seed was harvested and planted on GM agar containing 50µg/ml kanamycin. Plants surviving selection were transferred to soil and tested for complementation. Transformed plants and control Ws and *nim1* plants were sprayed with 300µm INA. Two days later, leaves were harvested for RNA extraction and PR-1 expression analysis. The plants were then sprayed with *Peronospora parasitica* (isolate EmWa) and grown as previously described. Ten days following fungal infection, plants were evaluated and scored positive or negative for fungal growth. All of the 15 transformed plants, as well as the Ws controls, were negative for fungal growth following INA treatment, while the *nim1* controls were positive for fungal growth. RNA was extracted and analyzed as described above for these transformants and controls. Ws controls and all 15 transformants showed PR-1 gene induction following INA treatment, while the *nim1* controls did not show PR-1 induction by INA.

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4. Isolation of a *NIM1* cDNA.

An Arabidopsis cDNA library made in the IYES expression vector (Elledge et al, 1991, PNAS 88, 1731-1735) was plated and plaque lifts were performed. Filters were hybridized with a ³²P-labeled PCR product generated from the gene region containing nim-1. 14 positives were identified from a screen of approximately 150,000 plaques. Each plaque was purified and plasmid DNA was recovered. cDNA inserts were digested out of the vector using EcoRI, agarose-gel-purified and sequenced. Sequence obtained from the longest cDNA is indicated in Figure 15. To confirm that we had obtained the 5' end of the cDNA, a Gibco BRL 5' RACE kit was used following manufacturer's instructions. The resulting RACE products were sequenced and found to include the additional bases indicated in Figure 15. The transcribed region present in both cDNA clones and detected in RACE is shown as capital letters in Figure 15. Changes in the alleles are shown above the DNA strand. Capitals indicate the presence of the sequence in a cDNA clone or detected after RACE PCR.

Example 9

Characterization of the *NIM1* gene

The multiple sequence alignment was constructed using Clustal V (Higgins, Desmond G. and Paul M. Sharp (1989), Fast and sensitive multiple sequence alignments on a microcomputer, CABIOS 5:151-153) as part of the DNA* (1228 South Park Street, Madison Wisconsin, 53715) Lasergene Biocomputing Software package for the Macintosh (1994).

It has been determined that certain regions of the *NIM1* protein are homologous in amino acid sequence to 4 different rice cDNA protein products. The homologies were

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identified using the *NIM1* sequences in a GenBank BLAST search. Comparisons of the regions of homology in *NIM1* and the rice cDNA products are shown in Figure 19. The *NIM1* protein fragments show from 36 to 48% identical amino acid sequences with the 4 rice products.

Example 10

Phenotypic characterization of the various *nim1* alleles

1. Analysis of chemical responsiveness in *nim1* alleles.

We analyzed the differences among the various *nim1* alleles in terms of chemical induction of PR gene expression and *Peronospora parasitica* resistance (see Figures 17 and 18).

Mutant plants were treated with chemical inducers and then assayed for PR gene expression and disease resistance.

2. Plant growth and chemical application.

Wild-type seeds and seeds for each of the *nim1* alleles (*nim1-1*, -2, -3, -4, -5, -6) were sown onto MetroMix 300 growing media, covered with a transparent plastic dome and placed at 4° C in the dark for 3 days. After 3 days of 4° C treatment the plants were moved to a phytotron for 2 weeks. At approximately 2 weeks post-planting, germinated seedlings had produced 4 true leaves. Plants were then treated with H₂O, 5mM SA, 300 uM BTH or 300 uM INA. Chemicals were applied as a fine mist to completely cover the seedlings using a chromister. Water control plants were returned to the growing phytotron while the chemically treated plants were held in a separate but identical phytotron. After 3 days plants were divided into 2 groups. One group was harvested for RNA extraction and analysis. The second group was inoculated with *P. parasitica*.

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3. *Peronospora parasitica* inoculation and analysis.

P. parasitica isolate 'EmWa' is a *P.p.* isolate that is compatible in the Ws ecotype. Compatible isolates are those that are capable of causing disease on a particular host. The *P. parasitica* isolate 'NoCo' is incompatible on Ws but compatible on the Columbia ecotype. Incompatible pathogens are recognized by the potential host, eliciting a host response that prevents disease development. At 3 days post-chemical application water and chemically treated plants were inoculated with the compatible 'EmWa' isolate. 'NoCo' inoculation was conducted on water treated plants only. Following inoculation plants were covered with a clear plastic dome to maintain high humidity required for successful *P. parasitica* infection and placed in a growing chamber with 19° C day/17° C night temperatures and 8h light/16h dark cycles.

At various timepoints after inoculation plants were analyzed microscopically to assess symptom development. Under magnification sporulation of the fungus can be observed at very early stages of disease development. The percentage of plants/pot showing sporulation at 5d, 6d, 7d, 11d and 14d after inoculation was determined and the density of sporulation was also recorded.

Figure 18 shows the disease assessment of the various *nim1* alleles following *P. parasitica* inoculation. The most distinguishing timepoints are 5 and 6 days post-inoculation. At 5 days post-inoculation *nim1-4* shows ~80% infection under all inducing chemical treatments performed, clearly indicating that this allele/genotype has the most severe disease susceptibility. At 6 days post-inoculation, *nim1*, -2, -3, -4 and -6 show significant disease incidence under all inducing chemical treatments. However, *nim1-5* shows less infection than Ws wild-type under all treatments at day 6. Therefore, *nim1-5* is the most disease resistant of the various *nim1* alleles. *nim1-2* appears intermediate with respect to disease susceptibility after BTH but not the other inducing treatments.

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PR-1 gene expression indicates that *nim1-4* is the least responsive to all of the inducing chemicals tested (Figure 17), while *nim1-5* shows elevated levels of PR-1 expression in the absence of inducers. These PR-1 gene expression results are consistent with the disease assessment performed with *P. parasitica* (Figure 18) and indicate that *nim1* alleles can cause resistance or susceptibility.

The samples obtained above were used to analyze *NIM1* gene expression (Figure 17). In wildtype plants *NIM1* mRNA was present in the untreated control samples. Following treatment with SA, INA, BTH or infection with a compatible pathogen the *NIM1* mRNA accumulated to higher levels. Differences in *NIM1* message (mRNA) abundance were observed in the *nim1* alleles compared to wildtype. The abundance of *NIM1* mRNA in untreated mutant plants was lower than observed in the wildtype with the exception of *nim1-2* and *-5* where the amounts were similar. The *nim1-1*, *-3* and *-4* had low levels of *NIM1* message while the *nim1-6* had very low accumulation of *NIM1* mRNA. Increases in *NIM1* mRNA following SA, INA or BTH were observed in *nim1-1*, *-2*, *-3* but not *nim1-5* or *-6*. However, this increase was less than observed in wildtype plants. Following pathogen infection additional bands hybridizing to the *NIM1* cDNA probe were observed in both wildtype and mutants and the *NIM1* mRNA level was elevated relative to untreated controls, except in *nim1-6*.

Figure 18 shows the disease resistance assessment via infection rating of the various *nim1* alleles as well as the NahG plants at various times after inoculation with *Peronospora parasitica*. WsWT indicates the Ws wild type parent line in which the *nim1* alleles are found. The various *nim1* alleles are indicated in the table and the NahG plant is indicated also. The NahG plant has been previously published. (Delaney et al. Science 266, pp. 1247-1250 (1994)). The NahG *Arabidopsis* is also described in WO 95/19443.

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The *NahG* gene is a gene from *Pseudomonas putida* that converts salicylic acid to catechol, thereby eliminating the accumulation of salicylic acid, a necessary signal transduction component for SAR in plants. Thus, *NahG Arabidopsis* plants do not display normal SAR. In addition, they show much greater susceptibility in general to pathogens. Therefore, the *NahG* plants serve as a kind of universal susceptibility control. In addition, the *NahG* plants still respond to the chemical inducers INA and BTH; this is shown in the bottom two panels of the Figure 17.

From Figure 18 it can be seen that the *nim1-4* and *nim1-6* alleles seem to be the most severe; this is most easily observable at the earlier time points, described earlier in the results section herein, and from the results set forth in the EmWa BTH panel, the lowest panel, in the Figure. In addition, the *nim1-5* allele shows the greatest response to both INA and BTH and therefore it is the weakest *nim1* allele.

The *NahG* plants show very good response to both INA and BTH and look very similar to the *nim1-5* allele. However, at late time points, Day 11 in the Figure, the disease resistance induced in the *NahG* plants begins to fade, and there is a profound difference between INA and BTH in that the INA-induced resistance fades much faster and more severely than the resistance induced in the *NahG* plants by BTH. Also seen in these experiments is that INA and BTH induced very good resistance in Ws to EmWa, and the *nim1-1*, *nim1-2* and other *nim1* alleles show virtually no response to SA or INA in regard to disease resistance.

Figure 18 lists the percent of plants that are showing sporulation after infection with the EmWa race of *P. parasitica*, and each of the bar graphs indicates the number of days after infection that the disease resistance was rated.

Northern analysis analyzing the expression of the SAR gene PR1 was also performed on the same samples, as shown in Figure 17. Figure 17 shows that the wild type

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plant shows very good response to salicylate, INA, BTH and also to pathogen infection, as manifested by enhanced PR1 gene expression. The *nim1-1* allele, on the other hand, shows only very limited response to all the chemical inducers including pathogen.

The pathogen induction is at least several fold lower in the *nim1-1* allele than it is in the wild type. The *nim1-2*, *nim1-3* and *nim1-6* alleles show response similar to the *nim1-1* allele to the various treatments. However, the *nim1-4* allele shows virtually no expression in response to any of the inducers used. Basically, background level is all that is observed. The *nim1-5* allele shows a very high background level relative to controls with water and that background level is maintained in all the treatments; however, there is limited or no induction by the chemical inducers.

The NahG plants serve as a good control, showing that they are unable to induce PR-1 in the presence of SA; on the other hand, INA and BTH both induce very strong high level expression of PR-1. The effect of pathogen infection is similar to that of SA; there is no expression of PR-1 in the EmWa-treated NahG plants.

These same RNA samples produced in the induction studies were also probed with a *NIM1* gene using a full-length cDNA clone as probed. In Figure 16 it can be seen that INA induces the *NIM1* gene in the wild type Ws allele. However, the *nim1-1* mutation allele shows a lower basal level expression of the *NIM1* gene, and it is not inducible by INA. This is similar to what is observed in the *nim1-3* allele and the *nim1-6* allele. The *nim1-2* allele shows approximately normal levels in the untreated sample and shows similar induction to that of the wild type sample, as does the *nim1-4* allele. The *nim1-5* allele seems to show higher basal level expression of the *NIM1* gene and much stronger expression when induced by chemical inducers.

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The induction of *NIM1* by chemical inducers of resistance and other inducers is consistent with its role in pathogen defense and is also further evidence that we have obtained the right gene in the 9.9 kb region.

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SEQUENCE LISTING

(1) GENERAL INFORMATION:

(i) APPLICANT:

- (A) NAME: Novartis AG
- (B) STREET: Schwarzwaldallee 215
- (C) CITY: Basel
- (E) COUNTRY: Switzerland
- (F) POSTAL CODE (ZIP): 4002
- (G) TELEPHONE: +41 61 69 11 11
- (H) TELEFAX: + 41 61 696 79 76
- (I) TELEX: 962 991

(ii) TITLE OF INVENTION: GENE CONFERRING DISEASE RESISTANCE IN
PLANTS AND USES THEREOF

(iii) NUMBER OF SEQUENCES: 11

(v) COMPUTER READABLE FORM:

- (A) MEDIUM TYPE: Floppy disk
- (B) COMPUTER: IBM PC compatible
- (C) OPERATING SYSTEM: PC-DOS/MS-DOS
- (D) SOFTWARE: PatentIn Release #1.0, Version #1.30

(2) INFORMATION FOR SEQ ID NO:1:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 9919 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (genomic)

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

TGATCATGAA TTGCGTGTAG GGTGTGTTTT TAAAGATAGG GATGAGCTGA AGAAGGCGGT	60
GGACTGGTGT TCCATTAGAG GGCAGCAAAA GTGTGTAGTA CAAGAGATTG AGAAGGACGA	120
GTATACGTTT AAATGCATCA GATGGAAATG CAATTGGTCG CGTCGGGCAG ATTGAATAGA	180
AGAACATGGA CTTGTTAAGA TAACTAAGTG TAGTTGGTCC ACATACTTGT TGTTCTATTA	240
AGCCGGAAAA CTTCAACTTG TAATTTCAG CAGAAGAGAT TGAGTGTCTG ATCAGGGTAC	300
AACCCACTCT AACAGCAGAG TTGAAAAGTT TGGTGACATG CTTAAAACCTT CAAAGCTGCG	360
GGCAGCAGAA CAGGAAGTAA TCAAAGATCA GAGTTTCAGA GTATTGCCTA AACTAATTGG	420
CTGCATTTC AATCTAAT GGGCTACTTG TGGACTGCAA TATGAGCTTT TCCCTAATCC	480
TGAATTTGCA TCCTTCGGTG GCGCGTTTTG GCGGTTTCCA CAGTCCATTG AAGGGTTTCA	540
ACACTGTAGA CCTCTGATCA TAGTGGATTC AAAAGACTTG AACGGCAAGT ACCCTATGAA	600
ATTGATGATT TCCTCAGGAC TCGACGCTGA TGATTGCTTT TTCCCGCTTG CCTTCCGCT	660
TACCAAAGAA GTGTCCACTG ATAGTTGGCG TTGGTTTCTC ACTAATATCA GAGAGAAGGT	720
AACACAAAGG AAAGACGTTT GCCTCGTCTC CAGTCCTCAC CCGACATAG TTGCTGTTAT	780
TAACGAACCC GGATCACTGT GGCAAGAACC TTGGGTCTAT CACAGGTTCT GTCTGGATTG	840
TTTTTGCTTA CAATTCCATG ATATTTTGG AGACTACAAC CTGGTGAGCC TTGTGAAGCA	900
GGCTGGATCC ACAAGTCAGA AGGAAGAATT TGATTCCTAC ATAAAGGACA TCAAAAAGAA	960
GGACTCAGAA GCTCGGAAAT GGTTAGCCCA ATTCCCTCAA AATCAGTGGG CTCTGGCTCA	1020
TGACCACTGG TCGGAGATAT GGAGTCATGA CGATAGAAAC AGAAGATTG AGGGCAATTT	1080
GTGAAAGCTT TCAGTCTCTT GGTCTATCAG TGACAGCGAA CGCACCTGCA CATGTGGGAA	1140

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GTTC AATCG AAGAAGTTTC CATGTATGCA CCCAGAAATG GTGCAAAGGA TTGTAACTT 1200

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TAAGAGGCAA GAGTCTCACC GACGACGATG AGAGAGTTTA CGGTTAGACC TCTTTCCACC 1740

GGTTGATTTC GATGTGGAAG AAGTCGAATC TGTCAGGGAC GAATTTCCCTA ATTCCAAATT 1800

GTCCCTCACTA AAGGCCTTCT TTAGTGCTC TTGTATTTCC ATGTACCTTT GCTTCTTTTG 1860

TAGTCGTTTC TCAGCAGTGT CGTCTTCTCC GCAAGCCAGT TGAGTCAAGT CCTCACAGTT 1920

CATAATCTGG TCGAGCACTG CCGAACAGCG CGGGAAGAAT CGTTTCCCGA GTTCCACTGA 1980

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CGCTTTTAGT CTACTTTGAT GCTCTCTAG GATTCTGAAA GGTGCTATCT TTACACCCGG 2160

TGATGTCTC TTCGTACCAG TGAGACGGTC AGGCTCGAGG CTAGTCACTA TGAACTCACA 2220

TGTTCCCTTC ATTTGGGCGA TCTCCATTGC AGCTTGTGCT TCCGTTGGAA AAAGACGTTG 2280

AGCAAGTGCA ACTAAACAGT GGACGACACA AAGAATAGTT ATCATTAGTT CACTCAGTTT 2340

CCTAATAGAG AGGACATAAA TTTAATTCAA ACATATAAGA AATAAGACTT GATAGATACC 2400

TCTATTTTCA AGATCGAGCA GCGTCATCTT CAATTCATCG GCCGCCACTG CAAAAGAGGG 2460

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GGCTTGTTTT GCGATCATGA GTGCGGTTCT ACCTTCCAAA GTTGCTTCTG ATGCACTTGC	2640
ACCTTTTTCC AATAGAGATA GTATCAATTG TGGCTCCTTC CGCATCGCAG CAACATGAAG	2700
CACCGTATAT CCCCTCGGAT TCCTATGGTT GACATCGGCA AGATCAAGTT TTAAAAGATC	2760
TGTTGCGGTC TTCACATTGC AATATGCAAC AGCGAAATGA AGAGCACACG CATCATCTAG	2820
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CTTATGTACA TTCGAGACAT GTTCTTTTAC TTTAGGTACC TCCAAACCAA GCTCTTTACG	2940
TCTATCAATT ATCTCTTTAA CAAGCTCTTC CGGCAATGAC TTTTCAAGAC TAACCATATC	3000
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ACATATATTA GCAAGCTTGA GTATAACCAA TGTGTCCTCT ATAACAACCT TGTCTACAAC	3120
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TAGCGGCGGC TAAAGCGCTC TTGAAGAAAG AGCTTCTCGC TGACAAAACG CACCGGTGGA	3540
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TGAGTACTTG TTCGGCGGCC AGATAAACAA TAGAGGAGTC GGTGTTATCG GTAGCGACGA	3720

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AACTAGTGCT GCTGATTTC TAAGAATCGG CGAATCCATC AATGGTGGTG TCCATCAACA	3780
GGTTCCGATG AATTGAAATT CACAAATTAA AGAGATCTCT GCTAATCAAC GAAGAGACCT	3840
TATCAACTGG ATTTGGTTAA AGATCGAAGA TAACCATTGA CGAGCAGAGC CAAGTCAAGT	3900
CAACGAGAGT GGTGGTGAGA TATGAAGAAG CATCCTCGTC CCACGGTTTA CATTTACCA	3960
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GCTAATTTTT GTATATGAGA AGTTCAATCC GGTTCCGTAA GCCCCTGAAC CAACTAGAT	4140
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AGCATGTTTA CACTATAATT AAATCAAGTC GAATATTTCC TGGAACTATT CTCCTGTTC	4440
TATAGCAAAT GAAAACGCTC TTCACAACAA AATCATTATA GATATAGGAA TAAATTACAT	4500
TAAAAACATG AAAGTCATAA TGAATATATT TTTTAAATTA GGATTTGATT TAAAAACAAT	4560
TATTGTATAC ATATAAAGA CTCTTTTAGT TATTTCCTT CAACTTCTCG TTCTGAATCA	4620
TGCGATAAAT CAGCTTTTTT AATAACTACG ACGTAAAAGC AAATTCATAA CACGTCTAAA	4680
CAAATTTGGC TCATCCTTCA CTTGATTGGT GTTTCCGGA CTCGATGTTG CTGGAACTG	4740
AGAAGAAGAA GGAATCTGCA TAATCACCTC TTGGTTCCTC ACCGGTAGAC TCATTTTGT	4800
GGATCGAAAA CGATCGAGAT CAGAAAATGA AAAGATAGGT TAAAGATGCC TATGAATACA	4860
ACAACGTAAG ATTATGTTGA ATAAACAGAG TACTTTATAT AGGAGTTATA ATAAGGTAAA	4920
TAAATATTG CTTCCGCGT TTTTACTTT TGTATTCTT AAATGATAAG TTAAATTAGG	4980
ATAAGATTG TATGATTTA AGTAAATTTA CAATACTCT CTATAACTCA ATAGCATCAC	5040

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TAAACCCAAT GACATATCCC AACCCCTTCAC TTCTGGCTTT GGTATGACCT GATACTGTTT 6480
AGTGGTTGGT TTGAAGACTA TGTATCCACG TGATGGTTTT GTATACTTAA CACAAAGCAA 6540
TATCCCATGA CTTGCATCAC AAGCTTCGAT CTTTATCATT CCGGGTGGCA GAAAGTCGAT 6600
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CGTGTTTGT CCATATATTT GTTGCTTATA AATATATTCA TATAACAATG TTTGCATTAA	7800
GCTTTTAAGA AGCACAAAAC CATATAACAA AATTAAATAT TCCTATCCCT ACCAAAAAAA	7860
AAAATTAAAT ATTCCCTACAG CCTTGTTGAT TATTTTATGC CCTACGTTGA GCCTTGTTGA	7920
CTAGTTTGCA TTTGTCGGTC CATTTCTTCT TCCGTCCAGA TCAACCCTCT CGTAATCAGA	7980
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TATCCAGTGA CGAATAACAC CTAGCTTCCC TTCGTAGCTG ACTAACTCTG GGAATAAACC	8280
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ACTCCTCACT TTTGGGTTTG GTATGATCTG ATACTGTTTT GTTGTGGTT TGCAGACTAT	8640
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GGTAGAGAAC CGACGATGAT GAATATACAA GTGTTTATAA GTATCACAAA TTGCCTTTTT	9120
CTTCGCTAGT CCCAAAACAA GCAAATTAAC CAAAGATAAA ATCTTCATTA ATGTTTTCCT	9180
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GGCGAAACGA ATATTCTGAT TCTAAAGATA GTAAAAATGA ATTTTGATGA AGGGAATACT	9840
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(2) INFORMATION FOR SEQ ID NO:2:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 5655 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

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(ii) MOLECULE TYPE: DNA (genomic)

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

(ix) FEATURE:

(A) NAME/KEY: exon

(B) LOCATION: 2787..3347

(D) OTHER INFORMATION: /product= "1st exon of NIM1"

(ix) FEATURE:

(A) NAME/KEY: exon

(B) LOCATION: 3427..4162

(D) OTHER INFORMATION: /product= "2nd exon of NIM1"

(ix) FEATURE:

(A) NAME/KEY: exon

(B) LOCATION: 4271..4474

(D) OTHER INFORMATION: /product= "3rd exon of NIM1"

(ix) FEATURE:

(A) NAME/KEY: exon

(B) LOCATION: 4586..4866

(D) OTHER INFORMATION: /product= "4th exon of NIM1"

(ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: join(2787..3347, 3427..4162, 4271..4474,
4586..4866)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

TGTGATGCAA GTCATGGGAT ATTGCTTTGT GTTAAGTATA CAAAACCATC ACGTGGATAC	60
ATAGTCTTCA AACCAACCAC TAAACAGTAT CAGGTCATAC CAAAGCCAGA AGTGAAGGGT	120
TGGGATATGT CATTGGGTTT AGCGGTAATC GGATTGAACC CTTTCCGGTA TAAAATACAA	180
AGGCTTTCGC AGTCTCGGCG TATGTGTATG TCTCGGGGTA TCTACCATTT GAATCACAGA	240

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ACTTTTATGT GCGAAGTTTT CGATTCTGAT TCGTTTACCT GGAAGAGATT AGAAAAATTG	300
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CATTGGATAT CACGGAACAA CAATGTGATC CGGTTTTGTC TCAAAACCGA AACTTGGTCC	420
TTCTTCCATA CTCCGAACTC TGATGTTTTT TCAGGATTAG TCAGATACGA AGGGAAGCTA	480
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CTCTACAACA TAAACGCAGA GAAGTTGAAT TTAGTTTATG CAAAAAAGA GGGATCTGAT	720
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ATCTAGTGAT GTTTAATTGT TTTTATAAG GTAAAAAGGA ATATTGAATT TTGTTTCTTA	960
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TATTTTATTA TATCAAGGGT TCCTGTTTAT ACTTGAAAAC AGTTACTGTA TAGAAAATAG	1080
TGTCCCAATT TTCTCTCTTA AATAATATAT TAGTTAATAA AAGATATTTT AATATATTAG	1140
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CTATAATAGA TGGTAGAAGA TAAAAAATT ATATCAGATT GATTCAATTA AATTTTATAA	1440
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AACGCGGAAA GCAATAATTT ATTTACCTTA TTATAACTCC TATATAAAGT ACTCTGTTTA 1680
TTCAACATAA TCTTACGTTG TTGTATTCAT AGGCATCTTT AACCTATCTT TTCATTTTCT 1740
GATCTCGATC GTTTTCGATC CAACAAAATG AGTCTACCGG TGAGGAACCA AGAGGTGATT 1800
ATGCAGATTC CTTCTTCTTC TCAGTTTCCA GCAACATCGA GTCCGGAAAA CACCAATCAA 1860
GTGAAGGATG AGCCAAATTT GTTTAGACGT GTTATGAATT TGCTTTTACG TCGTAGTTAT 1920
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ATATACATTA CAAAACCTAT GTGAATAAAG CATGAACTT AATATACGTT CCCTTTATCA 2280
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TTAAAAATA ATATTTATAT ATTTATATGA AAATAACGAA CCGGATGAAA AATAAATTTT 2400
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TTTAACCAAA TCCAGTTGAT AAGGTCTCTT CGTTGATTAG CAGAGATCTC TTTAATTGT 2760
GAATTTCAAT TCATCGGAAC CTGTTG ATG GAC ACC ACC ATT GAT GGA TTC GCC 2813

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Met Asp Thr Thr Ile Asp Gly Phe Ala															
1								5							
GAT TCT TAT GAA ATC AGC AGC ACT AGT TTC GTC GCT ACC GAT AAC ACC	2861														
Asp Ser Tyr Glu Ile Ser Ser Thr Ser Phe Val Ala Thr Asp Asn Thr															
10	15				20				25						
GAC TCC TCT ATT GTT TAT CTG GCC GCC GAA CAA GTA CTC ACC GGA CCT	2909														
Asp Ser Ser Ile Val Tyr Leu Ala Ala Glu Gln Val Leu Thr Gly Pro															
	30				35				40						
GAT GTA TCT GCT CTG CAA TTG CTC TCC AAC AGC TTC GAA TCC GTC TTT	2957														
Asp Val Ser Ala Leu Gln Leu Leu Ser Asn Ser Phe Glu Ser Val Phe															
	45				50				55						
GAC TCG CCG GAT GAT TTC TAC AGC GAC GCT AAG CTT GTT CTC TCC GAC	3005														
Asp Ser Pro Asp Asp Phe Tyr Ser Asp Ala Lys Leu Val Leu Ser Asp															
	60				65				70						
GGC CGG GAA GTT TCT TTC CAC CGG TGC GTT TTG TCA GCG AGA AGC TCT	3053														
Gly Arg Glu Val Ser Phe His Arg Cys Val Leu Ser Ala Arg Ser Ser															
	75				80				85						
TTC TTC AAG AGC GCT TTA GCC GCC GCT AAG AAG GAG AAA GAC TCC AAC	3101														
Phe Phe Lys Ser Ala Leu Ala Ala Ala Lys Lys Glu Lys Asp Ser Asn															
	90				95				100				105		
AAC ACC GCC GCC GTG AAG CTC GAG CTT AAG GAG ATT GCC AAG GAT TAC	3149														
Asn Thr Ala Ala Val Lys Leu Glu Leu Lys Glu Ile Ala Lys Asp Tyr															
	110				115				120						
GAA GTC GGT TTC GAT TCG GTT GTG ACT GTT TTG GCT TAT GTT TAC AGC	3197														
Glu Val Gly Phe Asp Ser Val Val Thr Val Leu Ala Tyr Val Tyr Ser															
	125				130				135						
AGC AGA GTG AGA CCG CCG CCT AAA GGA GTT TCT GAA TGC GCA GAC GAG	3245														
Ser Arg Val Arg Pro Pro Pro Lys Gly Val Ser Glu Cys Ala Asp Glu															
	140				145				150						
AAT TGC TGC CAC GTG GCT TGC CGG CCG GCG GTG GAT TTC ATG TTG GAG	3293														
Asn Cys Cys His Val Ala Cys Arg Pro Ala Val Asp Phe Met Leu Glu															
	155				160				165						

SUBSTITUTE SHEET (RULE 26)

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GTT CTC TAT TTG GCT TTC ATC TTC AAG ATC CCT GAA TTA ATT ACT CTC	3341
Val Leu Tyr Leu Ala Phe Ile Phe Lys Ile Pro Glu Leu Ile Thr Leu	
170 175 180 185	
 TAT CAG GTAAACACC ATCTGCATTA AGCTATGGTT ACACATTCAT GAATATGTTC	3397
Tyr Gln	
 TTACTTGAGT ACTTGATTTT GTATTTTCAG AGG CAC TTA TTG GAC GTT GTA GAC	3450
Arg His Leu Leu Asp Val Val Asp	
190 195	
 AAA GTT GTT ATA GAG GAC ACA TTG GTT ATA CTC AAG CTT GCT AAT ATA	3498
Lys Val Val Ile Glu Asp Thr Leu Val Ile Leu Lys Leu Ala Asn Ile	
200 205 210	
 TGT GGT AAA GCT TGT ATG AAG CTA TTG GAT AGA TGT AAA GAG ATT ATT	3546
Cys Gly Lys Ala Cys Met Lys Leu Leu Asp Arg Cys Lys Glu Ile Ile	
215 220 225	
 GTC AAG TCT AAT GTA GAT ATG GTT AGT CTT GAA AAG TCA TTG CCG GAA	3594
Val Lys Ser Asn Val Asp Met Val Ser Leu Glu Lys Ser Leu Pro Glu	
230 235 240	
 GAG CTT GTT AAA GAG ATA ATT GAT AGA CGT AAA GAG CTT GGT TTG GAG	3642
Glu Leu Val Lys Glu Ile Ile Asp Arg Arg Lys Glu Leu Gly Leu Glu	
245 250 255	
 GTA CCT AAA GTA AAG AAA CAT GTC TCG AAT GTA CAT AAG GCA CTT GAC	3690
Val Pro Lys Val Lys Lys His Val Ser Asn Val His Lys Ala Leu Asp	
260 265 270 275	
 TCG GAT GAT ATT GAG TTA GTC AAG TTG CTT TTG AAA GAG GAT CAC ACC	3738
Ser Asp Asp Ile Glu Leu Val Lys Leu Leu Leu Lys Glu Asp His Thr	
280 285 290	
 AAT CTA GAT GAT GCG TGT GCT CTT CAT TTC GCT GTT GCA TAT TGC AAT	3786
Asn Leu Asp Asp Ala Cys Ala Leu His Phe Ala Val Ala Tyr Cys Asn	
295 300 305	
 GTG AAG ACC GCA ACA GAT CTT TTA AAA CTT GAT CTT GCC GAT GTC AAC	3834
Val Lys Thr Ala Thr Asp Leu Leu Lys Leu Asp Leu Ala Asp Val Asn	
310 315 320	

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CAT AGG AAT CCG AGG GGA TAT ACG GTG CTT CAT GTT GCT GCG ATG CGG	3882
His Arg Asn Pro Arg Gly Tyr Thr Val Leu His Val Ala Ala Met Arg	
325 330 335	
AAG GAG CCA CAA TTG ATA CTA TCT CTA TTG GAA AAA GGT GCA AGT GCA	3930
Lys Glu Pro Gln Leu Ile Leu Ser Leu Leu Glu Lys Gly Ala Ser Ala	
340 345 350 355	
TCA GAA GCA ACT TTG GAA GGT AGA ACC GCA CTC ATG ATC GCA AAA CAA	3978
Ser Glu Ala Thr Leu Glu Gly Arg Thr Ala Leu Met Ile Ala Lys Gln	
360 365 370	
GCC ACT ATG GCG GTT GAA TGT AAT AAT ATC CCG GAG CAA TGC AAG CAT	4026
Ala Thr Met Ala Val Glu Cys Asn Asn Ile Pro Glu Gln Cys Lys His	
375 380 385	
TCT CTC AAA GGC CGA CTA TGT GTA GAA ATA CTA GAG CAA GAA GAC AAA	4074
Ser Leu Lys Gly Arg Leu Cys Val Glu Ile Leu Glu Gln Glu Asp Lys	
390 395 400	
CGA GAA CAA ATT CCT AGA GAT GTT CCT CCC TCT TTT GCA GTG GCG GCC	4122
Arg Glu Gln Ile Pro Arg Asp Val Pro Pro Ser Phe Ala Val Ala Ala	
405 410 415	
GAT GAA TTG AAG ATG ACG CTG CTC GAT CTT GAA AAT AGA G	4162
Asp Glu Leu Lys Met Thr Leu Leu Asp Leu Glu Asn Arg	
420 425 430	
GTATCTATCA AGTCTTATTT CTTATATGTT TGAATTAAAT TTATGTCCTC TCTATTAGGA	4222
AACTGAGTGA ACTAATGATA ACTATTCTTT GTGTCGTCCA CTGTTTAG TT GCA CTT	4278
Val Ala Leu	
435	
GCT CAA CGT CTT TTT CCA ACG GAA GCA CAA GCT GCA ATG GAG ATC GCC	4326
Ala Gln Arg Leu Phe Pro Thr Glu Ala Gln Ala Ala Met Glu Ile Ala	
440 445 450	
GAA ATG AAG GGA ACA TGT GAG TTC ATA GTG ACT AGC CTC GAG CCT GAC	4374
Glu Met Lys Gly Thr Cys Glu Phe Ile Val Thr Ser Leu Glu Pro Asp	
455 460 465	

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CGT CTC ACT GGT ACG AAG AGA ACA TCA CCG GGT GTA AAG ATA GCA CCT	4422
Arg Leu Thr Gly Thr Lys Arg Thr Ser Pro Gly Val Lys Ile Ala Pro	
470 475 480	
TTC AGA ATC CTA GAA GAG CAT CAA AGT AGA CTA AAA GCG CTT TCT AAA	4470
Phe Arg Ile Leu Glu Glu His Gln Ser Arg Leu Lys Ala Leu Ser Lys	
485 490 495	
ACC G GTATGGATTC TCACCCACTT CATCGGACTC CTTATCACAA AAAACAAAAC	4524
Thr	
500	
TAAATGATCT TTAACATGG TTTTGTTACT TGCTGTCTGA CTTGTGTTTT TTTATCATCA	4584
G TG GAA CTC GGG AAA CGA TTC TTC CCG CGC TGT TCG GCA GTG CTC	4629
Val Glu Leu Gly Lys Arg Phe Phe Pro Arg Cys Ser Ala Val Leu	
505 510 515	
GAC CAG ATT ATG AAC TGT GAG GAC TTG ACT CAA CTG GCT TGC GGA GAA	4677
Asp Gln Ile Met Asn Cys Glu Asp Leu Thr Gln Leu Ala Cys Gly Glu	
520 525 530	
GAC GAC ACT GCT GAG AAA CGA CTA CAA AAG AAG CAA AGG TAC ATG GAA	4725
Asp Asp Thr Ala Glu Lys Arg Leu Gln Lys Lys Gln Arg Tyr Met Glu	
535 540 545	
ATA CAA GAG ACA CTA AAG AAG GCC TTT AGT GAG GAC AAT TTG GAA TTA	4773
Ile Gln Glu Thr Leu Lys Lys Ala Phe Ser Glu Asp Asn Leu Glu Leu	
550 555 560	
GGA AAT TCG TCC CTG ACA GAT TCG ACT TCT TCC ACA TCG AAA TCA ACC	4821
Gly Asn Ser Ser Leu Thr Asp Ser Thr Ser Ser Thr Ser Lys Ser Thr	
565 570 575	
GGT GGA AAG AGG TCT AAC CGT AAA CTC TCT CAT CGT CGT CGG TGA	4866
Gly Gly Lys Arg Ser Asn Arg Lys Leu Ser His Arg Arg Arg *	
580 585 590	
GACTCTTGCC TCTTAGTGTA ATTTTGCTG TACCATATAA TTCTGTTTTT ATGATGACTG	4926
TAACTGTTTA TGTCTATCGT TGGCGTCATA TAGTTTCGCT CTTCGTTTTG CATCCTGTGT	4986
ATTATTGCTG CAGGTGTGCT TCAAACAAAT GTTGTAACAA TTTGAACCAA TGGTATACAG	5046

SUBSTITUTE SHEET (RULE 26)

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ATTTGTAATA TATATTTATG TACATCAACA ATAACCCATG ATGGTGTTAC AGAGTTGCTA 5106
 GAATCAAAGT GTGAAATAAT GTCAAATTGT TCATCTGTTG GATATTTTCC ACCAAGAACC 5166
 AAAAGAATAT TCAAGTTCCC TGAAGTTCTG GCAACATTCA TGTTATATGT ATCTTCCTAA 5226
 TTCTTCCTTT AACCTTTTGT AACTCGAATT ACACAGCAAG TTAGTTTCAG GTCTAGAGAT 5286
 AAGAGAACAC TGAGTGGGCG TGTAAGGTGC ATTCTCCTAG TCAGCTCCAT TGCATCCAAC 5346
 ATTTGTGAAT GACACAAGTT AACAATCCTT TGCACCATTT CTGGGTGCAT ACATGGAAAC 5406
 TTCTTCGATT GAAACTTCCC ACATGTGCAG GTGCGTTCGC TGTCACTGAT AGACCAAGAG 5466
 ACTGAAAGCT TTCACAAATT GCCCTCAAAT CTTCTGTTTC TATCGTCATG ACTCCATATC 5526
 TCCGACCACT GGTCA TGAGC CAGAGCCCAC TGATTTTGAG GGAATTGGGC TAACCATTTC 5586
 CGAGCTTCTG AGTCCTTCTT TTTGATGTCC TTTATGTAGG AATCAAATTC TTCCTTCTGA 5646
 CTGTGGAT 5655

(2) INFORMATION FOR SEQ ID NO:3:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 594 amino acids
 (B) TYPE: amino acid
 (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

Met Asp Thr Thr Ile Asp Gly Phe Ala Asp Ser Tyr Glu Ile Ser Ser
 1 5 10 15
 Thr Ser Phe Val Ala Thr Asp Asn Thr Asp Ser Ser Ile Val Tyr Leu
 20 25 30
 Ala Ala Glu Gln Val Leu Thr Gly Pro Asp Val Ser Ala Leu Gln Leu
 35 40 45

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Leu	Ser	Asn	Ser	Phe	Glu	Ser	Val	Phe	Asp	Ser	Pro	Asp	Asp	Phe	Tyr	50	55	60	
Ser	Asp	Ala	Lys	Leu	Val	Leu	Ser	Asp	Gly	Arg	Glu	Val	Ser	Phe	His	65	70	75	80
Arg	Cys	Val	Leu	Ser	Ala	Arg	Ser	Ser	Phe	Phe	Lys	Ser	Ala	Leu	Ala	85	90	95	
Ala	Ala	Lys	Lys	Glu	Lys	Asp	Ser	Asn	Asn	Thr	Ala	Ala	Val	Lys	Leu	100	105	110	
Glu	Leu	Lys	Glu	Ile	Ala	Lys	Asp	Tyr	Glu	Val	Gly	Phe	Asp	Ser	Val	115	120	125	
Val	Thr	Val	Leu	Ala	Tyr	Val	Tyr	Ser	Ser	Arg	Val	Arg	Pro	Pro	Pro	130	135	140	
Lys	Gly	Val	Ser	Glu	Cys	Ala	Asp	Glu	Asn	Cys	Cys	His	Val	Ala	Cys	145	150	155	160
Arg	Pro	Ala	Val	Asp	Phe	Met	Leu	Glu	Val	Leu	Tyr	Leu	Ala	Phe	Ile	165	170	175	
Phe	Lys	Ile	Pro	Glu	Leu	Ile	Thr	Leu	Tyr	Gln	Arg	His	Leu	Leu	Asp	180	185	190	
Val	Val	Asp	Lys	Val	Val	Ile	Glu	Asp	Thr	Leu	Val	Ile	Leu	Lys	Leu	195	200	205	
Ala	Asn	Ile	Cys	Gly	Lys	Ala	Cys	Met	Lys	Leu	Leu	Asp	Arg	Cys	Lys	210	215	220	
Glu	Ile	Ile	Val	Lys	Ser	Asn	Val	Asp	Met	Val	Ser	Leu	Glu	Lys	Ser	225	230	235	240
Leu	Pro	Glu	Glu	Leu	Val	Lys	Glu	Ile	Ile	Asp	Arg	Arg	Lys	Glu	Leu	245	250	255	
Gly	Leu	Glu	Val	Pro	Lys	Val	Lys	Lys	His	Val	Ser	Asn	Val	His	Lys	260	265	270	

SUBSTITUTE SHEET (RULE 26)

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Ala Leu Asp Ser Asp Asp Ile Glu Leu Val Lys Leu Leu Leu Lys Glu
 275 280 285

Asp His Thr Asn Leu Asp Asp Ala Cys Ala Leu His Phe Ala Val Ala
 290 295 300

Tyr Cys Asn Val Lys Thr Ala Thr Asp Leu Leu Lys Leu Asp Leu Ala
 305 310 315 320

Asp Val Asn His Arg Asn Pro Arg Gly Tyr Thr Val Leu His Val Ala
 325 330 335

Ala Met Arg Lys Glu Pro Gln Leu Ile Leu Ser Leu Leu Glu Lys Gly
 340 345 350

Ala Ser Ala Ser Glu Ala Thr Leu Glu Gly Arg Thr Ala Leu Met Ile
 355 360 365

Ala Lys Gln Ala Thr Met Ala Val Glu Cys Asn Asn Ile Pro Glu Gln
 370 375 380

Cys Lys His Ser Leu Lys Gly Arg Leu Cys Val Glu Ile Leu Glu Gln
 385 390 395 400

Glu Asp Lys Arg Glu Gln Ile Pro Arg Asp Val Pro Pro Ser Phe Ala
 405 410 415

Val Ala Ala Asp Glu Leu Lys Met Thr Leu Leu Asp Leu Glu Asn Arg
 420 425 430

Val Ala Leu Ala Gln Arg Leu Phe Pro Thr Glu Ala Gln Ala Ala Met
 435 440 445

Glu Ile Ala Glu Met Lys Gly Thr Cys Glu Phe Ile Val Thr Ser Leu
 450 455 460

Glu Pro Asp Arg Leu Thr Gly Thr Lys Arg Thr Ser Pro Gly Val Lys
 465 470 475 480

Ile Ala Pro Phe Arg Ile Leu Glu Glu His Gln Ser Arg Leu Lys Ala
 485 490 495

Leu Ser Lys Thr Val Glu Leu Gly Lys Arg Phe Phe Pro Arg Cys Ser

500 505 510

Arg *

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 41 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS: not relevant
(D) TOPOLOGY: not relevant

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

Val His Tyr Ala Val Gln His Cys Asn
35 40

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(2) INFORMATION FOR SEQ ID NO:5:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 38 amino acids
- (B) TYPE: amino acid
- (C) STRANDEDNESS: not relevant
- (D) TOPOLOGY: not relevant

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

Pro	Thr	Gly	Lys	Thr	Ala	Leu	His	Leu	Ala	Ala	Glu	Met	Val	Ser	Pro
1				5				10					15		
Asp	Met	Val	Ser	Val	Leu	Leu	Asp	His	His	Ala	Asp	Xaa	Asn	Phe	Arg
				20				25					30		
Thr	Xaa	Asp	Gly	Val	Thr										
				35											

(2) INFORMATION FOR SEQ ID NO:6:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 41 amino acids
- (B) TYPE: amino acid
- (C) STRANDEDNESS: not relevant
- (D) TOPOLOGY: not relevant

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

Ile	Arg	Arg	Met	Arg	Arg	Ala	Leu	Asp	Ala	Ala	Asp	Ile	Glu	Leu	Val
1				5				10					15		

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Lys Leu Met Val Met Gly Glu Gly Leu Asp Leu Asp Asp Ala Leu Ala
20 25 30

Val His Tyr Ala Val Gln His Cys Asn
35 40

(2) INFORMATION FOR SEQ ID NO:7:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 27 amino acids
- (B) TYPE: amino acid
- (C) STRANDEDNESS: not relevant
- (D) TOPOLOGY: not relevant

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

Arg Arg Pro Asp Ser Lys Thr Ala Leu His Leu Ala Ala Glu Met Val
1 5 10 15

Ser Pro Asp Met Val Ser Val Leu Leu Asp Gln
20 25

(2) INFORMATION FOR SEQ ID NO:8:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 41 amino acids
- (B) TYPE: amino acid
- (C) STRANDEDNESS: not relevant
- (D) TOPOLOGY: not relevant

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

Ile Arg Arg Met Arg Arg Ala Leu Asp Ala Ala Asp Ile Glu Leu Val

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1 5 10 15
Lys Leu Met Val Met Gly Glu Gly Leu Asp Leu Asp Asp Ala Leu Ala
 20 25 30
Val His Tyr Ala Val Gln His Cys Asn
 35 40

(2) INFORMATION FOR SEQ ID NO:9:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 27 amino acids
- (B) TYPE: amino acid
- (C) STRANDEDNESS: not relevant
- (D) TOPOLOGY: not relevant

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

Arg Arg Pro Asp Ser Lys Thr Ala Leu His Leu Ala Ala Glu Met Val
1 5 10 15
Ser Pro Asp Met Val Ser Val Leu Leu Asp Gln
 20 25

(2) INFORMATION FOR SEQ ID NO:10:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 41 amino acids
- (B) TYPE: amino acid
- (C) STRANDEDNESS: not relevant
- (D) TOPOLOGY: not relevant

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:

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Ile Arg Arg Met Arg Arg Ala Leu Asp Ala Ala Asp Ile Glu Leu Val
 1 5 10 15

Lys Leu Met Val Met Gly Glu Gly Leu Asp Leu Asp Asp Ala Leu Ala
 20 25 30

Val His Tyr Ala Val Gln His Cys Asn
 35 40

(2) INFORMATION FOR SEQ ID NO:11:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 19 amino acids
- (B) TYPE: amino acid
- (C) STRANDEDNESS: not relevant
- (D) TOPOLOGY: not relevant

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

Pro Thr Gly Lys Thr Ala Leu His Leu Ala Ala Glu Met Val Ser Pro
 1 5 10 15

Asp Met Val

Applicant's or agent's file reference number	Pat. No. 21212/A/CGC 1909	International application No.	PCT/EP 97 / 01218
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INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

(PCT Rule 13bis)

A. The indications made below relate to the microorganism referred to in the description on page _____, line _____	
B. IDENTIFICATION OF DEPOSIT Further deposits are identified on an additional sheet <input type="checkbox"/>	
Name of depositary institution American Type Culture Collection (ATCC)	
Address of depositary institution (including postal code and country) 12301 Parklawn Drive Rockville, MD 20852 USA	
Date of deposit 25 September 1996 (25.09.96)	Accession Number 97736
C. ADDITIONAL INDICATIONS (leave blank if not applicable) This information is continued on an additional sheet <input type="checkbox"/>	
We request the Expert Solution where available	
D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designated States)	
E. SEPARATE FURNISHING OF INDICATIONS (leave blank if not applicable)	
The indications listed below will be submitted to the International Bureau later (specify the general nature of the indications e.g., "Accession Number of Deposit")	
For receiving Office use only	For International Bureau use only
<input checked="" type="checkbox"/> This sheet was received with the international application	<input type="checkbox"/> This sheet was received by the International Bureau on:
Authorized officer Y. Marinus-v.d. Nouweland	Authorized officer



American Type Culture Collection

11361 Parklawn Drive • Rockville, MD 20852 USA • Telephone: (301)231-6520 Telex: 998-768 ATCCROVE • FAX: 301-816-4346

BUDAPEST TREATY ON THE INTERNATIONAL RECOGNITION OF THE DEPOSIT OF MICROORGANISMS FOR THE PURPOSES OF PATENT PROCEDURE

INTERNATIONAL FORM

RECEIPT IN THE CASE OF AN ORIGINAL DEPOSIT ISSUED PURSUANT TO RULE 7.3
AND VIABILITY STATEMENT ISSUED PURSUANT TO RULE 10.2

To: (Name and Address of Depositor or Attorney)

Ciba-Geigy Corporation
Attention: Leslie B. Friedrich
P.O. Box 12257
Research Triangle Park, NC 27709

Deposited on Behalf of: Ciba-Geigy Corporation

Identification Reference by Depositor:

ATCC Designation

Coamid, D7

97736

The deposit was accompanied by: ☐ a scientific description ☐ a proposed taxonomic description indicated above.

The deposit was received September 25, 1996 by this International Depository Authority and has been accepted.

AT YOUR REQUEST:

☒ We will inform you of requests for the strain for 30 years.

The strain will be made available if a patent office signatory to the Budapest Treaty certifies one's right to receive, or if a U.S. Patent is issued citing the strain, and ATCC is instructed by the United States Patent & Trademark Office or the depositor to release said strain.

If the culture should die or be destroyed during the effective term of the deposit, it shall be your responsibility to replace it with living culture of the same.

The strain will be maintained for a period of at least 30 years after the date of deposit, and for a period of at least five years after the most recent request for a sample. The United States and many other countries are signatory to the Budapest Treaty.

The viability of the culture cited above was tested October 3, 1996. On that date, the culture was viable.

International Depository Authority: American Type Culture Collection, Rockville, Md. 20852 USA

Signature of person having authority to represent ATCC:

Barbara M. Halley
Barbara M. Halley, Administrator, ATCC Patent Depository

Date: October 7, 1996

cc: Andrea C. Walsh, Ph.D.

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Applicant's or agent's file reference number	PP, -21212/A/CGC 1909	International application	F T/EP 97 / 0 1 2 1 8
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INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

(PCT Rule 13bis)

A. The indications made below relate to the microorganism referred to in the description on page _____, line _____	
B. IDENTIFICATION OF DEPOSIT Further deposits are identified on an additional sheet <input type="checkbox"/>	
Name of depositary institution American Type Culture Collection (ATCC)	
Address of depositary institution (including postal code and country) 12301 Parklawn Drive Rockville, MD 20852 USA	
Date of deposit 13 June 1996 (13.06.96)	Accession Number 97606
C. ADDITIONAL INDICATIONS (leave blank if not applicable) This information is continued on an additional sheet <input type="checkbox"/>	
We request the Expert Solution where available	
D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designated States)	
E. SEPARATE FURNISHING OF INDICATIONS (leave blank if not applicable)	
The indications listed below will be submitted to the International Bureau later (specify the general nature of the indications e.g., "Accession Number of Deposit")	
For receiving Office use only	For International Bureau use only
<input checked="" type="checkbox"/> This sheet was received with the international application Y. Marinus-v.d. Nouweiland	<input type="checkbox"/> This sheet was received by the International Bureau on:
Authorized officer	Authorized officer

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American Type Culture Collection

13301 Parklawn Drive • Rockville, MD 20852 USA • Telephone: (301)231-5530 Telex: 898-055 ATCCNORTH • FAX: (301)770-2587

BUDAPEST TREATY ON THE INTERNATIONAL RECOGNITION OF THE DEPOSIT OF MICROORGANISMS FOR THE PURPOSES OF PATENT PROCEDURE

INTERNATIONAL FORM

RECEIPT IN THE CASE OF AN ORIGINAL DEPOSIT ISSUED PURSUANT TO RULE 7.3 AND VIABILITY STATEMENT ISSUED PURSUANT TO RULE 10.2

To: (Name and Address of Depositor or Attorney)

Ciba-Geigy Corporation
Attn: Leslie B. Friedrich
P.O. Box 12257
Research Triangle Park, NC 27709

RECEIVED

JUN 27 1996

CIBA-GEIGY
ABRU PATENT DEPT

Deposited on Behalf of: Ciba-Geigy Corporation

Identification Reference by Depositor:

ATCC Designation

Plasmid P1-18

97808

The deposit was accompanied by: ☐ a scientific description ☐ a proposed taxonomic description indicated above.

The deposit was received June 13, 1996 by this International Depositary Authority and has been accepted.

AT YOUR REQUEST: ☒ We will inform you of requests for the strain for 30 years.

The strain will be made available if a patent office signatory to the Budapest Treaty certifies one's right to receive, or if a U.S. Patent is issued citing the strain, and ATCC is instructed by the United States Patent & Trademark Office or the depositor to release said strain.

If the culture should die or be destroyed during the effective term of the deposit, it shall be your responsibility to replace it with living culture of the same.

The strain will be maintained for a period of at least 30 years from date of deposit, or five years after the most recent request for a sample, whichever is longer. The United States and many other countries are signatory to the Budapest Treaty.

The viability of the culture cited above was tested June 20, 1996. On that date, the culture was viable.

International Depositary Authority: American Type Culture Collection, Rockville, Md. 20852 USA

Signature of person having authority to represent ATCC:

Barbara M. Hailey
Barbara M. Hailey, Administrator, Patent Depository

Date: June 21, 1996

cc: Andrea C. Walsh, Ph.D.

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Applicant's or agent's file reference number	P -21212/A/CGC 1909	International application No PCT/EP 97, 01218
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INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

(PCT Rule 13bis)

A. The indications made below relate to the microorganism referred to in the description on page _____, line _____	
B. IDENTIFICATION OF DEPOSIT Further deposits are identified on an additional sheet <input type="checkbox"/>	
Name of depositary institution American Type Culture Collection (ATCC)	
Address of depositary institution (including postal code and country) 12301 Parklawn Drive Rockville, MD 20852 USA	
Date of deposit 08 May 1996 (08.05.96)	Accession Number 97543
C. ADDITIONAL INDICATIONS (leave blank if not applicable) This information is continued on an additional sheet <input type="checkbox"/>	
We request the Expert Solution where available	
D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designated States)	
E. SEPARATE FURNISHING OF INDICATIONS (leave blank if not applicable) The indications listed below will be submitted to the International Bureau later (specify the general nature of the indications e.g., "Accession Number of Deposit")	

For receiving Office use only	For International Bureau use only
<input checked="" type="checkbox"/> This sheet was received with the international application	<input type="checkbox"/> This sheet was received by the International Bureau on:
Authorized officer Y. Marinus-v.d. Nouweland	Authorized officer

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American Type Culture Collection

12301 Parklawn Drive • Rockville, MD 20852 USA • Telephone: (301)351-5320 Telex: 699-655 ATCCNORTH • FAX: 301-770-3587

BUDAPEST TREATY ON THE INTERNATIONAL RECOGNITION OF
THE DEPOSIT OF MICROORGANISMS FOR THE PURPOSES OF PATENT PROCEDURE

INTERNATIONAL FORM

RECEIPT IN THE CASE OF AN ORIGINAL DEPOSIT ISSUED PURSUANT TO RULE 7.3
AND VIABILITY STATEMENT ISSUED PURSUANT TO RULE 10.2

To: (Name and Address of Depositor or Attorney)

Ciba-Geigy Corporation
Attn: Leslie B. Friedrich
P.O. Box 12267
Research Triangle Park, NC 27709

RECEIVED

MAY 28 1996

CIBA-GEIGY
ABRU PATENT DEPT.

Deposited on Behalf of: Ciba-Geigy Corporation

Identification Reference by Depositor:

ATCC Designation

Plasmid, BAC4

97543

The deposit was accompanied by: ☐ a scientific description ☐ a proposed taxonomic description indicated above.

The deposit was received May 8, 1996 by this International Depository Authority and has been accepted.

AT YOUR REQUEST: ☒ We will inform you of requests for the strain for 30 years.

The strain will be made available if a patent office signatory to the Budapest Treaty certifies one's right to receive, or if a U.S. Patent is issued citing the strain, and ATCC is instructed by the United States Patent & Trademark Office or the depositor to release said strain.

If the culture should die or be destroyed during the effective term of the deposit, it shall be your responsibility to replace it with living culture of the same.

The strain will be maintained for a period of at least 30 years from date of deposit, or five years after the most recent request for a sample, whichever is longer. The United States and many other countries are signatory to the Budapest Treaty.

The viability of the culture cited above was tested May 17, 1996. On that date, the culture was viable.

International Depository Authority: American Type Culture Collection, Rockville, Md. 20852 USA

Signature of person having authority to represent ATCC:

Barbara M. Halley

Barbara M. Halley, Administrator, Patent Depository

Date: May 20, 1996

cc: Andree C. Walsh, Ph.D.

SUBSTITUTE SHEET (RULE 26)

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What Is Claimed Is:

1. An isolated DNA molecule comprising a *NIM1* gene.
2. An isolated DNA molecule according to claim 1, comprising the nucleotide sequence set forth in SEQ ID NO:2.
3. An isolated DNA molecule of about 9.9kb which encodes the *NIM1* gene product.
4. An isolated DNA molecule according to claim 1, comprising the nucleotide sequence set forth in SEQ ID NO:1.
5. An isolated DNA molecule of claim 1, encoding the aminoacid sequence of the *NIM1* gene product set forth in SEQ ID NO:2.
6. An isolated DNA molecule comprising a mutant gene of *NIM1* of claim 1, which is a *nim1* gene.
7. Clone BAC-04, ATCC Deposit No. 97543.
8. A chimeric gene comprising a promotor active in plant operably linked to a heterologuous DNA molecule encoding the aminoacid sequence of a *NIM1* gene product.
9. A chimeric gene comprising a promotor active in plant operably linked to the heterologuous DNA fragment according to claim 3.
10. A chimeric gene comprising a promotor active in plant operably linked to a heterologuous DNA molecule encoding the aminoacid sequence set forth in SEQ ID NO:2.
11. A chimeric gene comprising a promotor active in plant operably linked to a heterologuous DNA molecule encoding the aminoacid sequence of a *nim1* gene product.
12. A recombinant vector comprising the chimeric gene of anyone of claims 8 to 11.
13. A recombinant vector according to claim 12, wherein said vector is capable of being stably transformed into a host cell.
14. A recombinant vector according to claim 12, wherein said vector is capable of being stably transformed into a plant, plant seeds, plant tissue or plant cell.

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26. Use of an isolated DNA molecule according to claim 1 and variants thereof in a screening method for identifying compounds capable of inducing broad spectrum disease resistance in plants.

27. Use of a plant phenotyp according to claim 17 to identify an isolated gene fragment which allows expression of broad spectrum of disease resistance in plants.

28. Use of an isolated DNA molecule according to anyone of claims 1 to 5 to confer disease resistance to plant cells, plants and the progeny thereof.

29. Use of an isolated DNA molecule according to claim 6 to confer universal disease susceptibility to plant cells, plants and the progeny thereof.

30. Use of resistant plants and the progeny thereof according to claim 20 to incorporate the resistant trait into plant lines through breeding.

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15. A plant expression cassette comprising a chimeric gene of anyone of claims 8 to 11.
16. A plant expression cassette comprising a chimeric gene of claim 8 to 10.
17. A plant expression cassette comprising a chimeric gene of claim 11.
18. A plant expression cassette according to claim 15 to 17 which expresses the chimeric gene continuously or constitutively.
19. A plant, plant cells and the progeny thereof comprising the chimeric gene of anyone of claims 8 to 11.
20. A plant, plant cells and the progeny thereof comprising the chimeric gene of anyone of claims 8 to 10, which have a broad spectrum of disease resistance.
21. A plant, plant cells and the progeny thereof comprising the chimeric gene of claim 11.
22. A plant, plant cells and the progeny thereof of claim 19, wherein said plant is selected from the group consisting of gymnosperms, monocots, and dicots.
23. A plant, plant cells and the progeny thereof of claim 19, wherein said plant is a crop plant.
24. A plant, plant cells and the progeny thereof of claim 23, wherein said plant is selected from the group consisting of rice, wheat, barley, rye, corn, potato, carrot, sweet potato, sugar beet, bean, pea, chicory, lettuce, cabbage, cauliflower, broccoli, turnip, radish, spinach, asparagus, onion, garlic, eggplant, pepper, celery, carrot, squash, pumpkin, zucchini, cucumber, apple, pear, quince, melon, plum, cherry, peach, nectarine, apricot, strawberry, grape, raspberry, blackberry, pineapple, avocado, papaya, mango, banana, soybean, tobacco, tomato, sorghum and sugarcane.
25. Use of an isolated DNA molecule according to claim 1, gene to confer disease resistance in plants.

Fig. 1

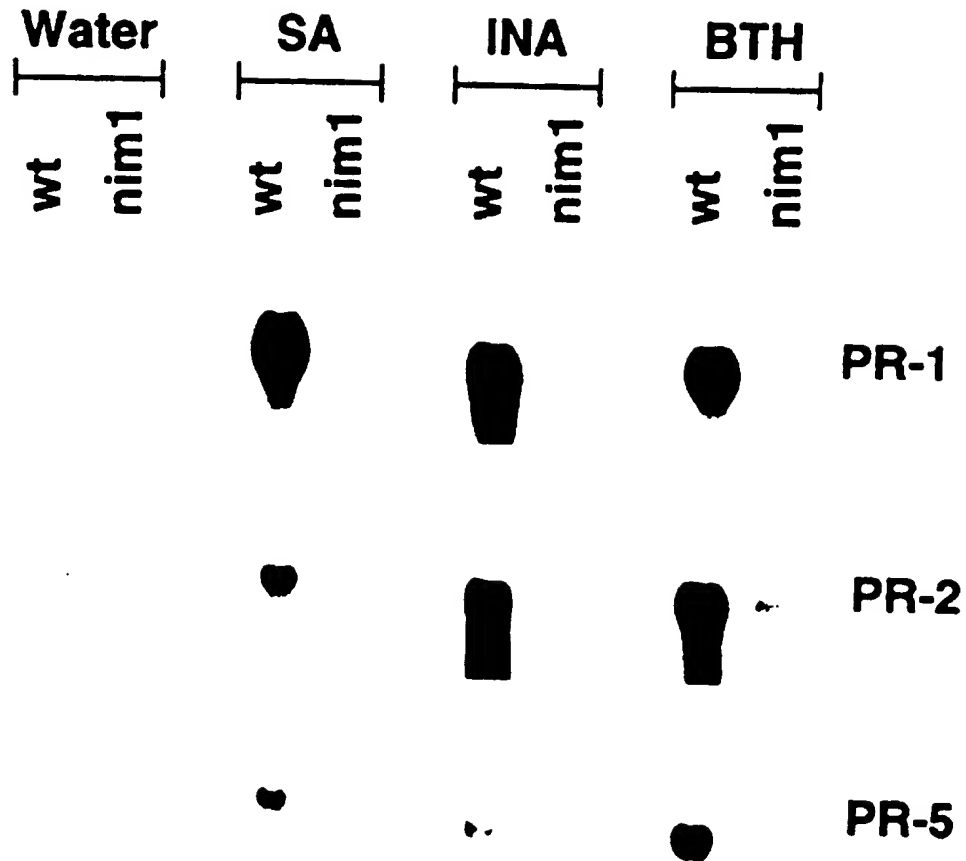


Fig. 2

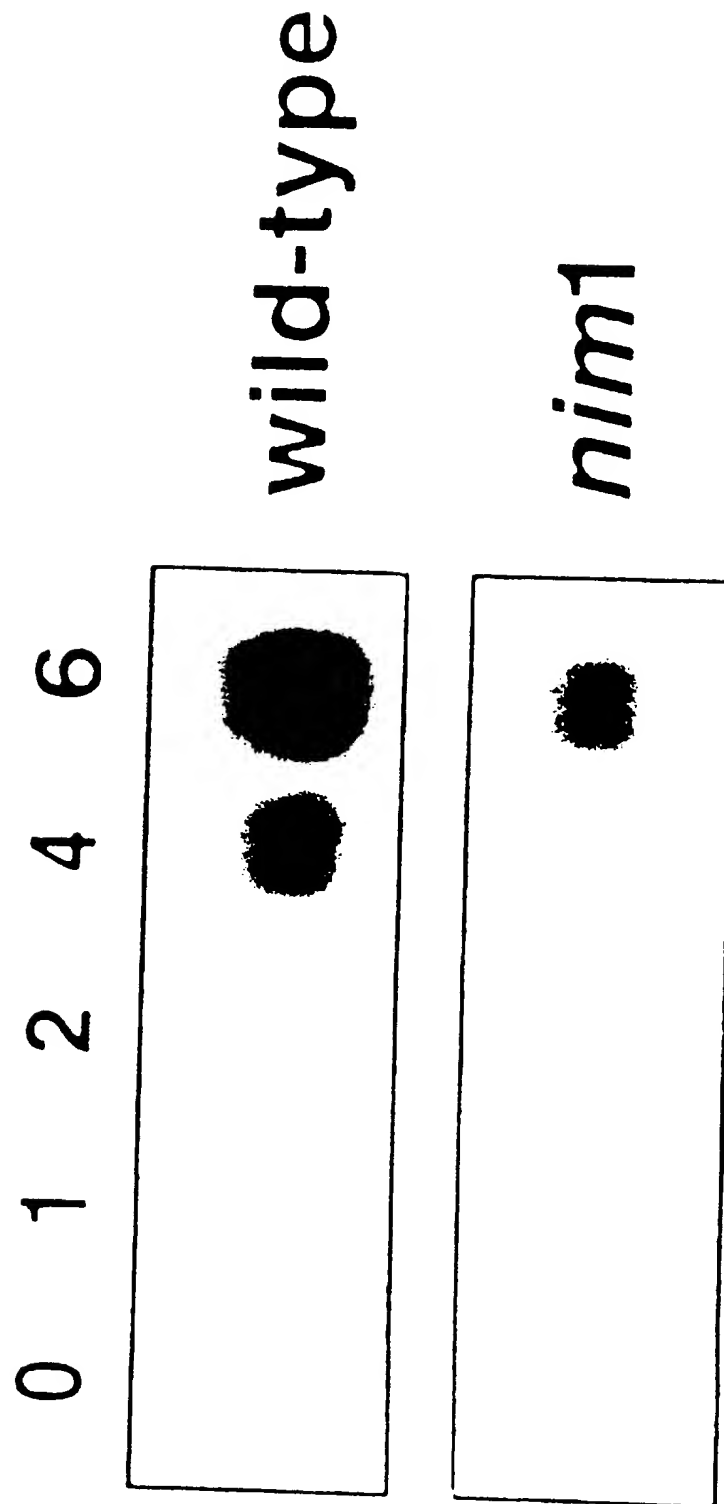
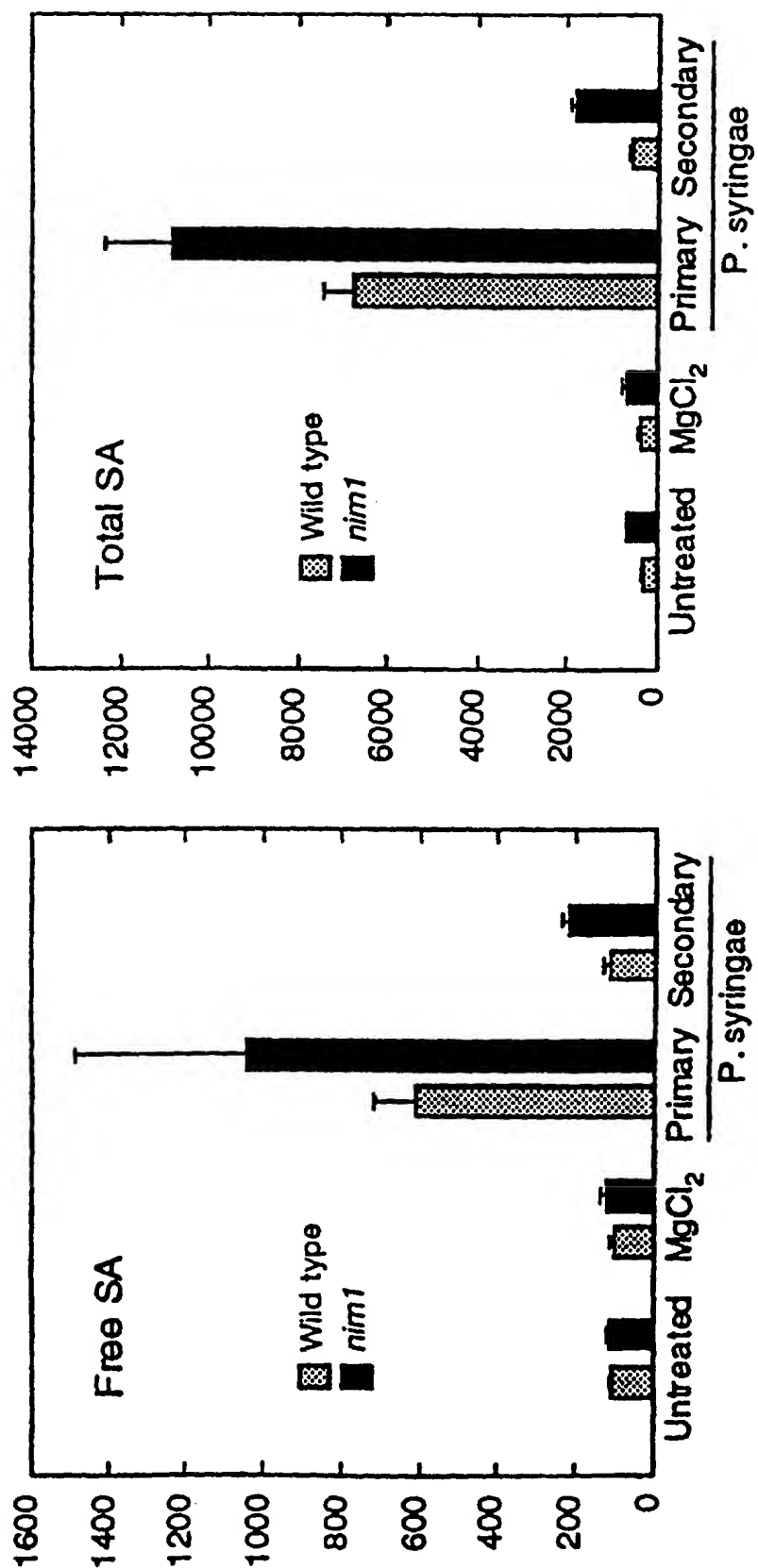


Fig. 3



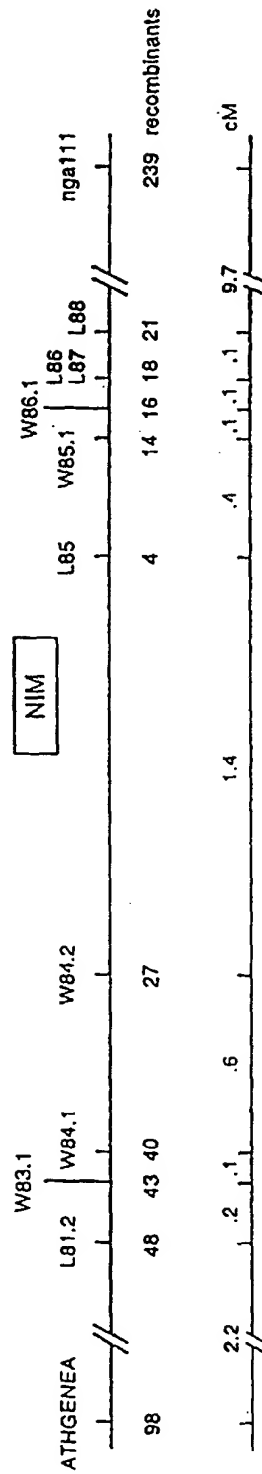
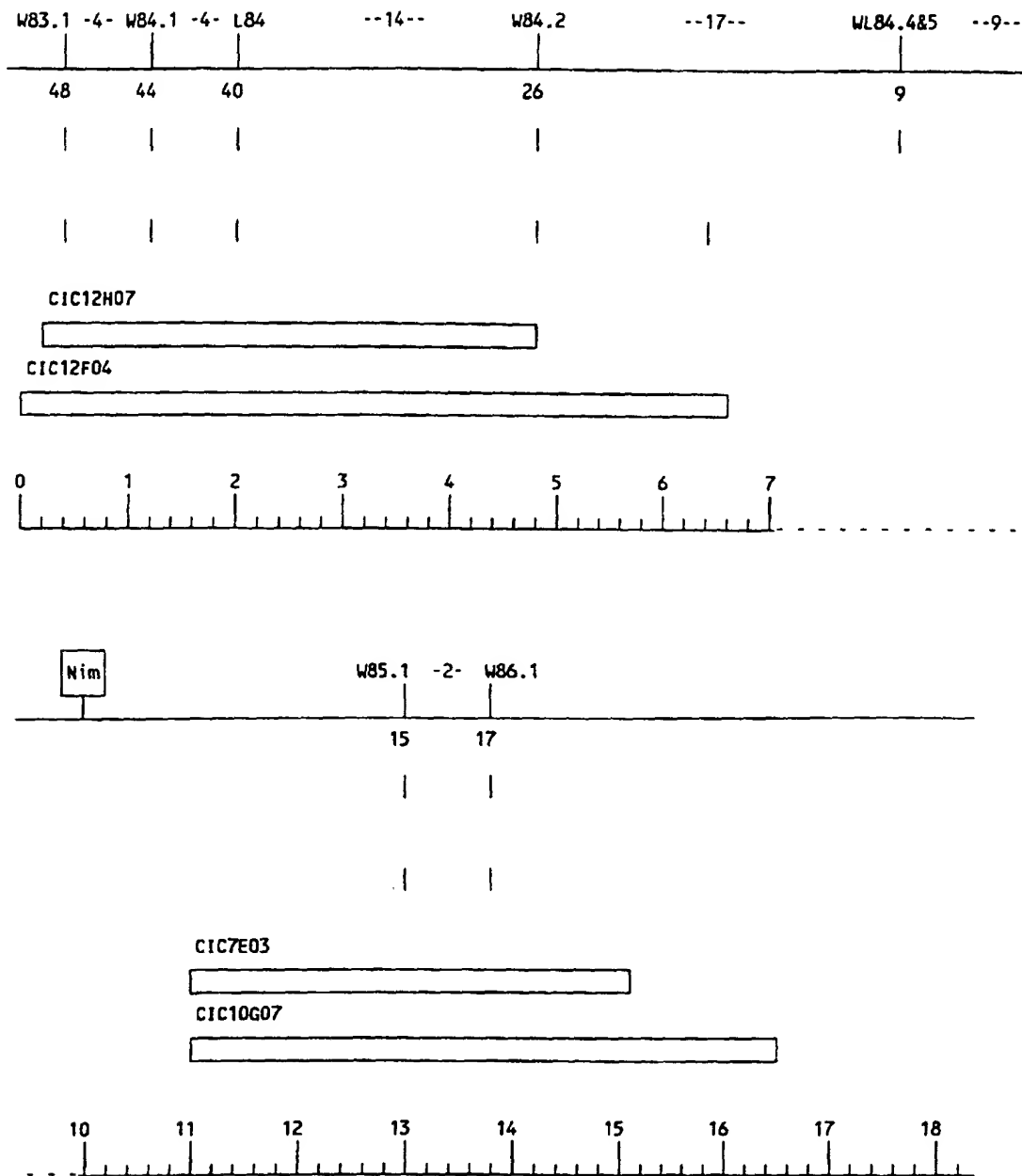


Figure 4. Genetic map of the NIM region, showing the approximate position of NIM, positions of the markers, the number of recombinants that are identified by each marker, and the genetic distances (cM) between the markers.

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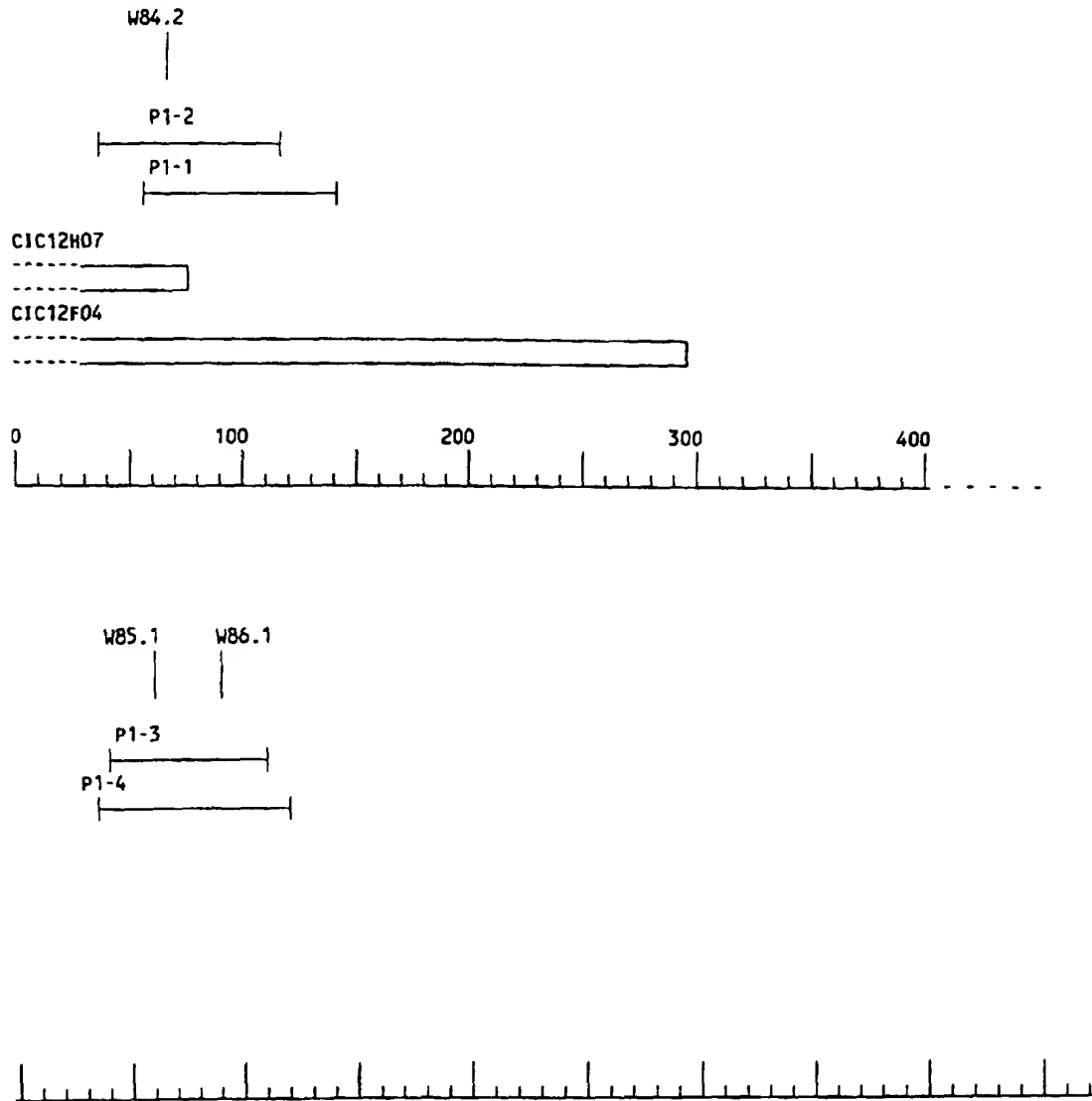
fig. 5



Schematic representation of the Nim flanking YACs with regard to the genetic map and the AFLP markers. Below the physical distance is indicated in n x 100 kb. Only the positions of markers W84.2 and WL84.4&5 were determined very accurately, the other markers can only be positioned within a certain interval.

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Fig. 6



Positions of P1 and BAC clones with respect to the flanking AFLP markers and YACs. The positions of clones P1-3 and P1-4 with respect to YACs 10G07 and 7E03 were not determined. At the bottom the physical distance in kb's is indicated. The physical distance and extent of overlaps are best estimates and not exact values.

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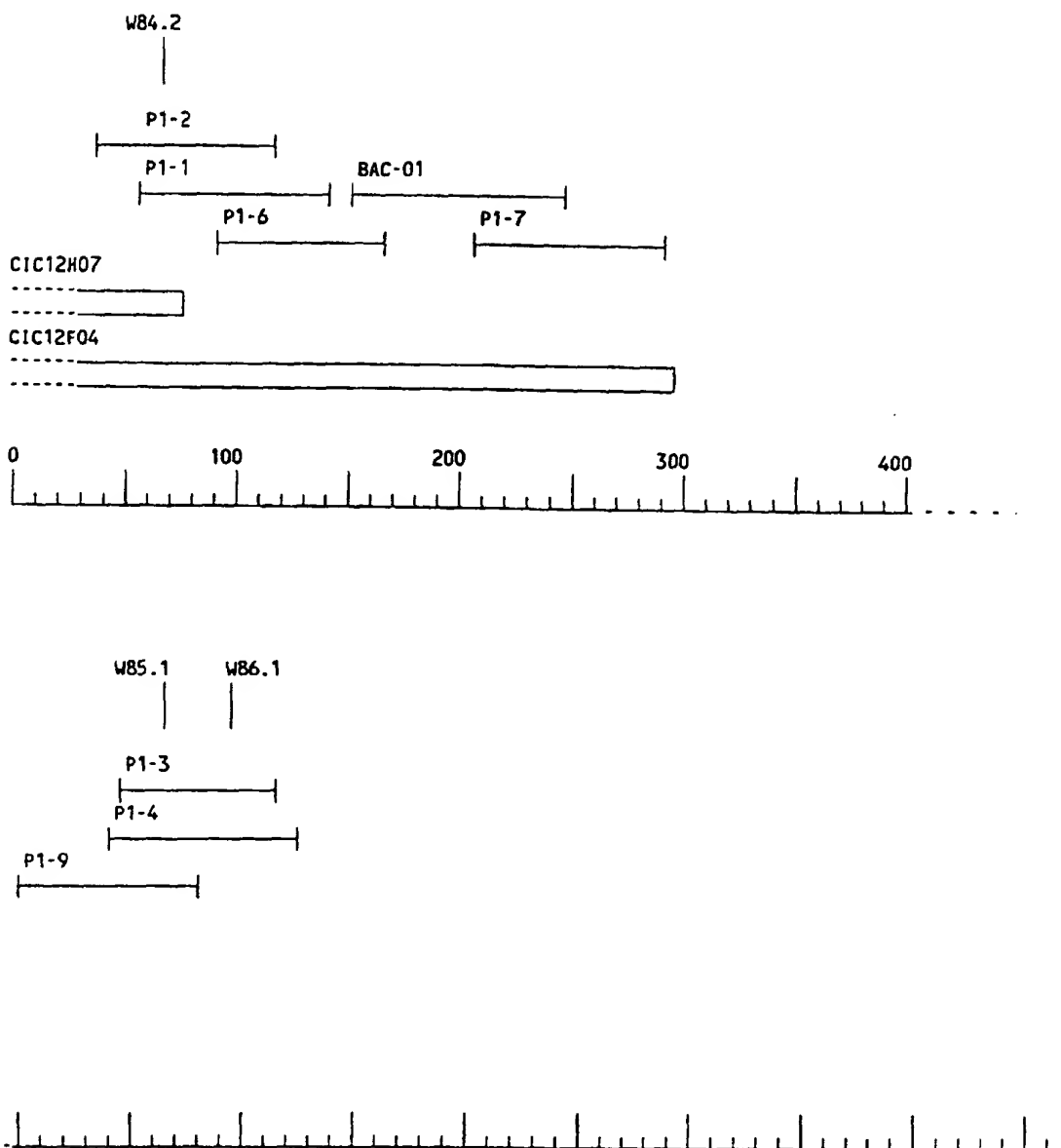


Fig. 7

Extended P1 and BAC contig covering the South end of YAC CIC12F04 and the flanking markers. Clones P1-3, P1-4 and P1-9 overlapped completely with YACs 10G07 and 7E03 and were not positioned with respect to these YACs. At the bottom the physical distance in kb's is indicated. The physical distance and extent of overlaps are best estimates and not exact values.

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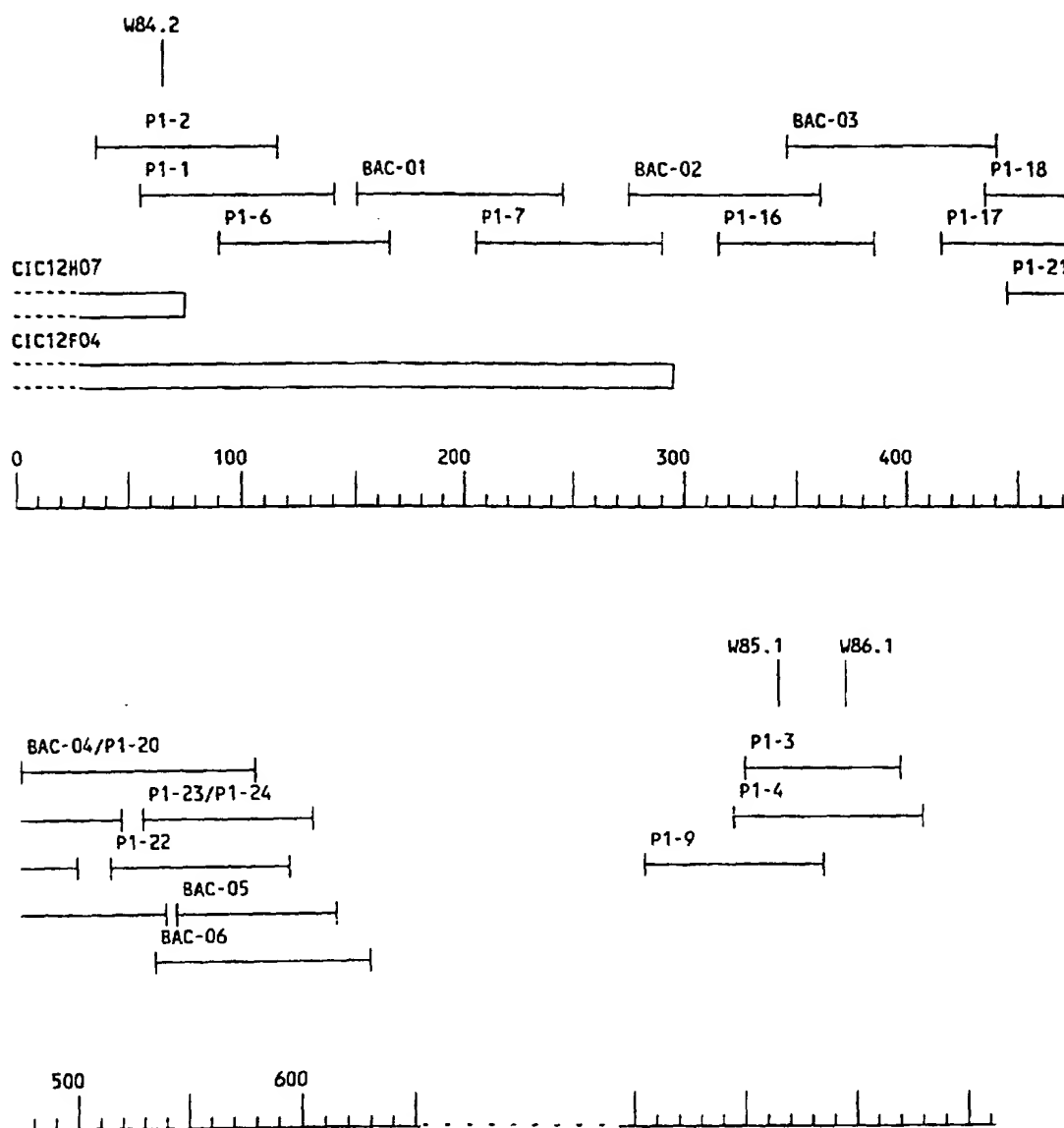


Fig. 8

Schematic representation of all identified P1 and BAC clones and their relative positions in the Nim region. At the bottom the physical distance in kb's is indicated. The physical distance and extent of overlaps are best estimates and not exact values.

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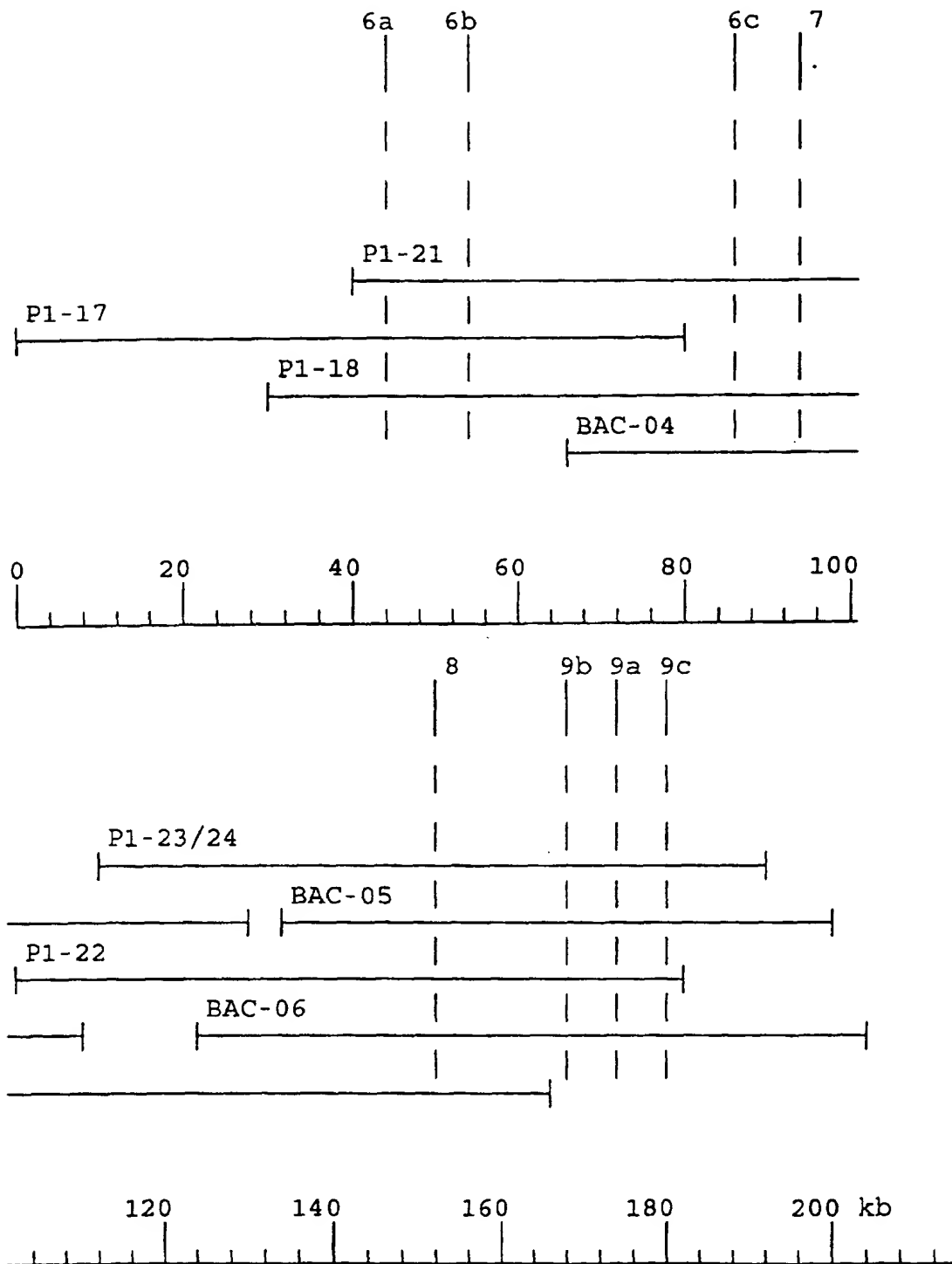


Fig. 9

Integrated genetic and physical fine map of the NIM region. The scale on the bottom is in kb.

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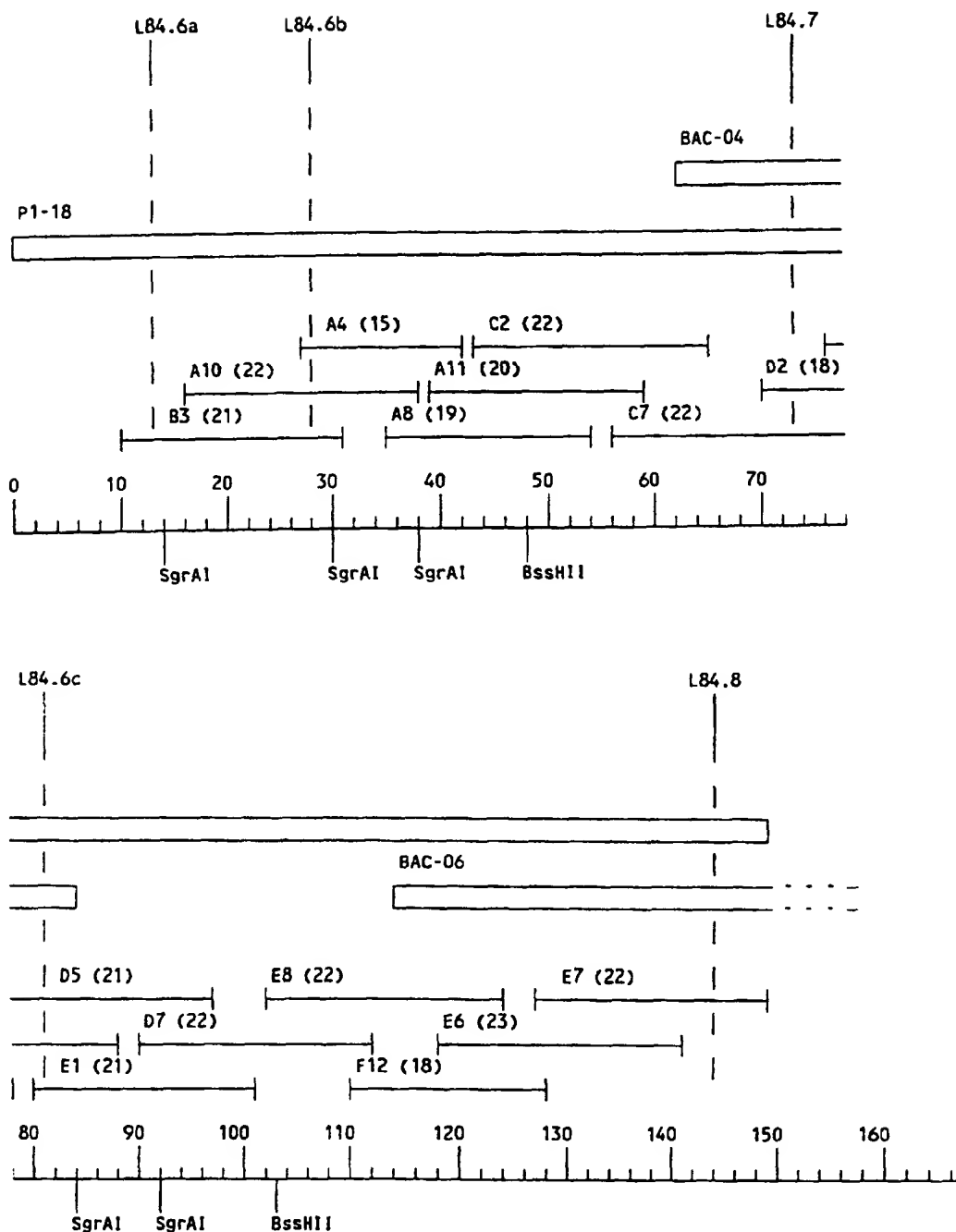


Fig. 10

Integrated map of the NIM region. This map is a best fit of the AFLP fingerprinting and restriction mapping data. Shown are the relevant AFLP markers, the BAC and P1 clones, the cosmid contig, the sizes of the cosmid inserts (in parentheses) and some restriction sites. The scale on the bottom is in kb.

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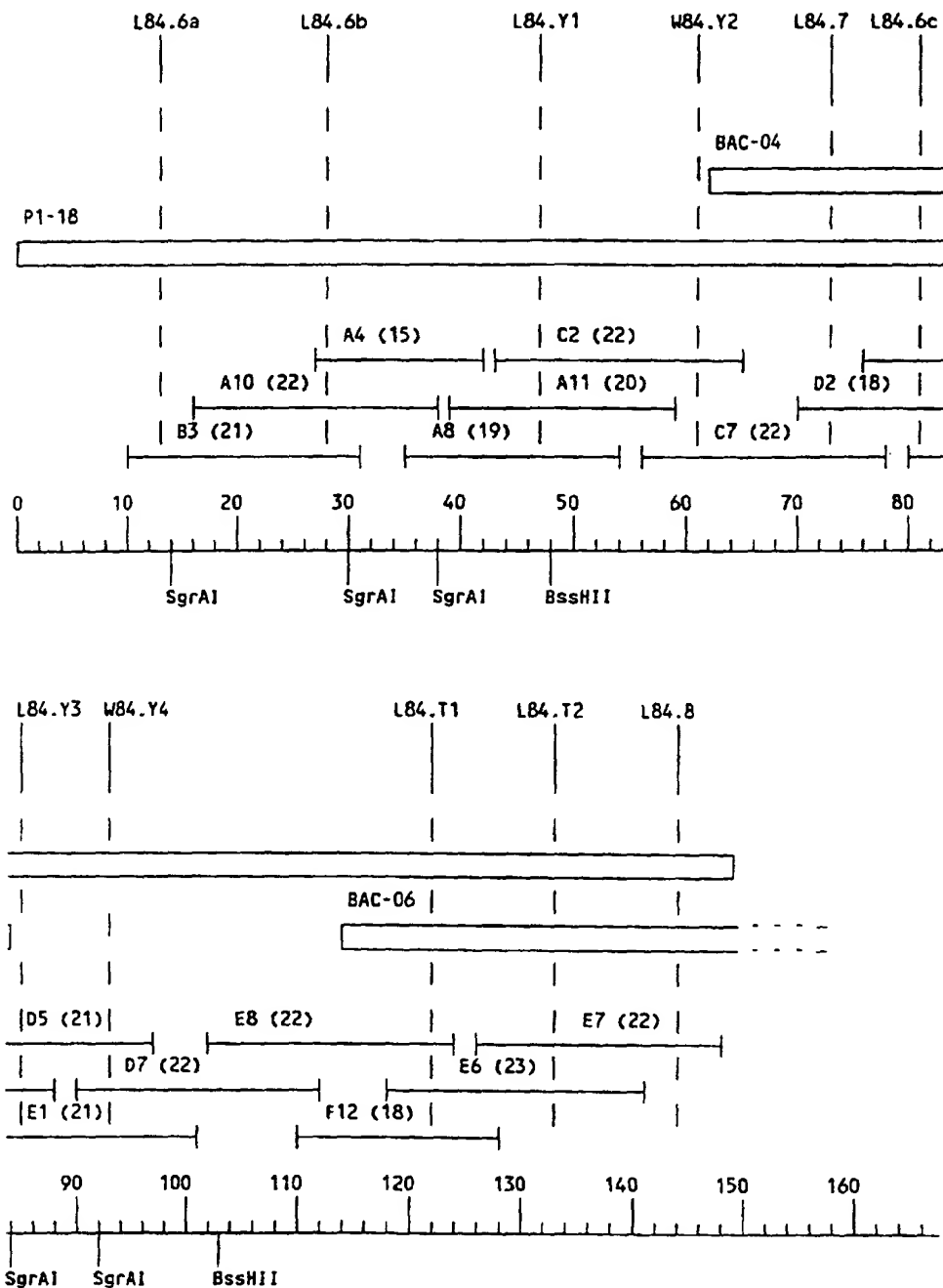


Fig. 11

Integrated map of the NIM region including the new AFLP markers. This map is a best fit of the AFLP fingerprinting and restriction mapping data. Shown are the relevant AFLP markers, the BAC and P1 clones, the cosmid contig, the sizes of the cosmid inserts (in parentheses) and some restriction sites. The scale on the bottom is in kb.

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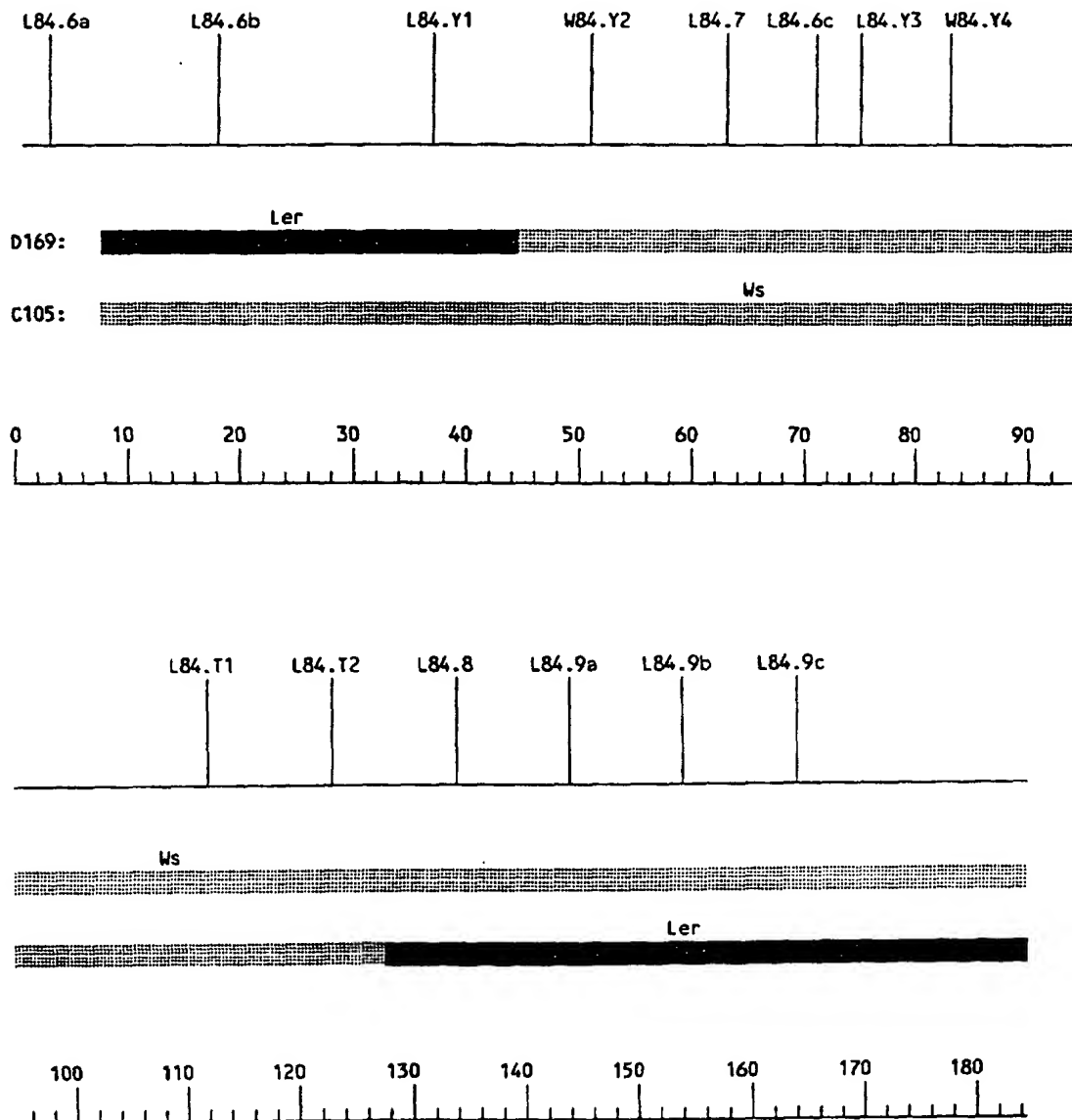
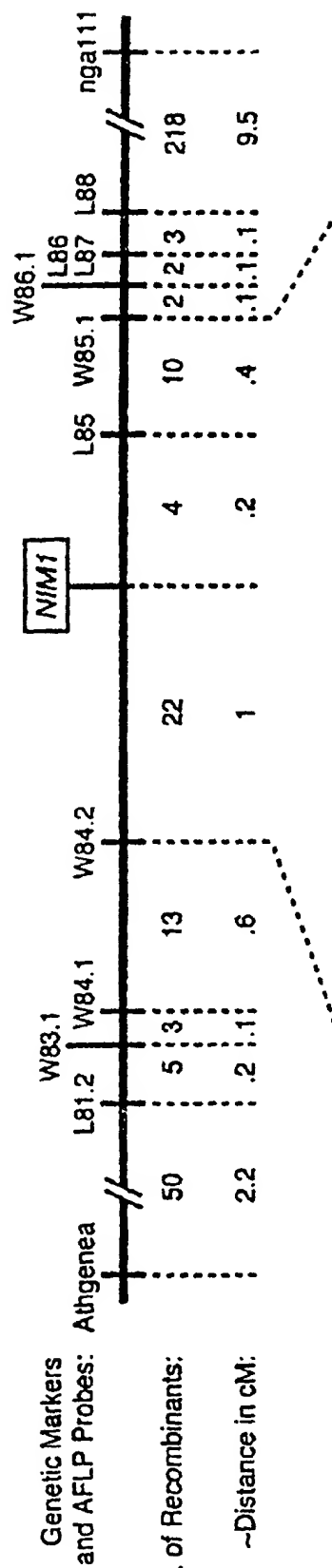
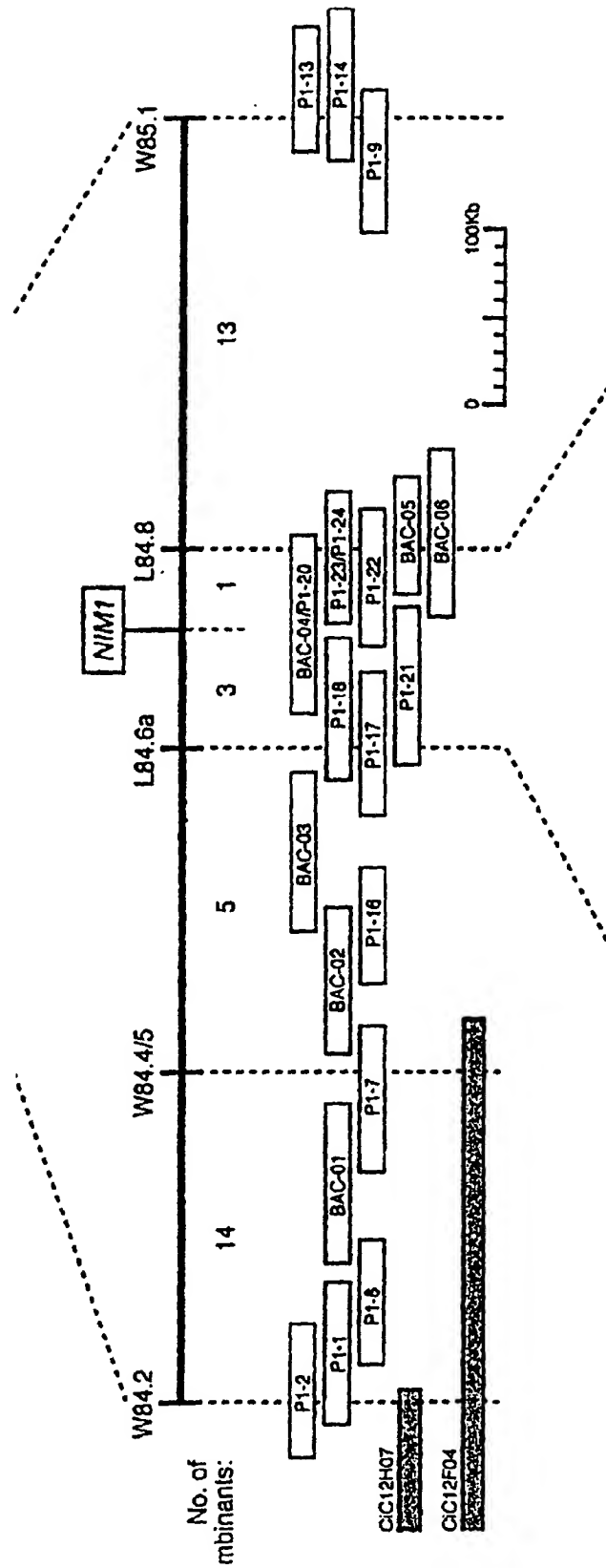


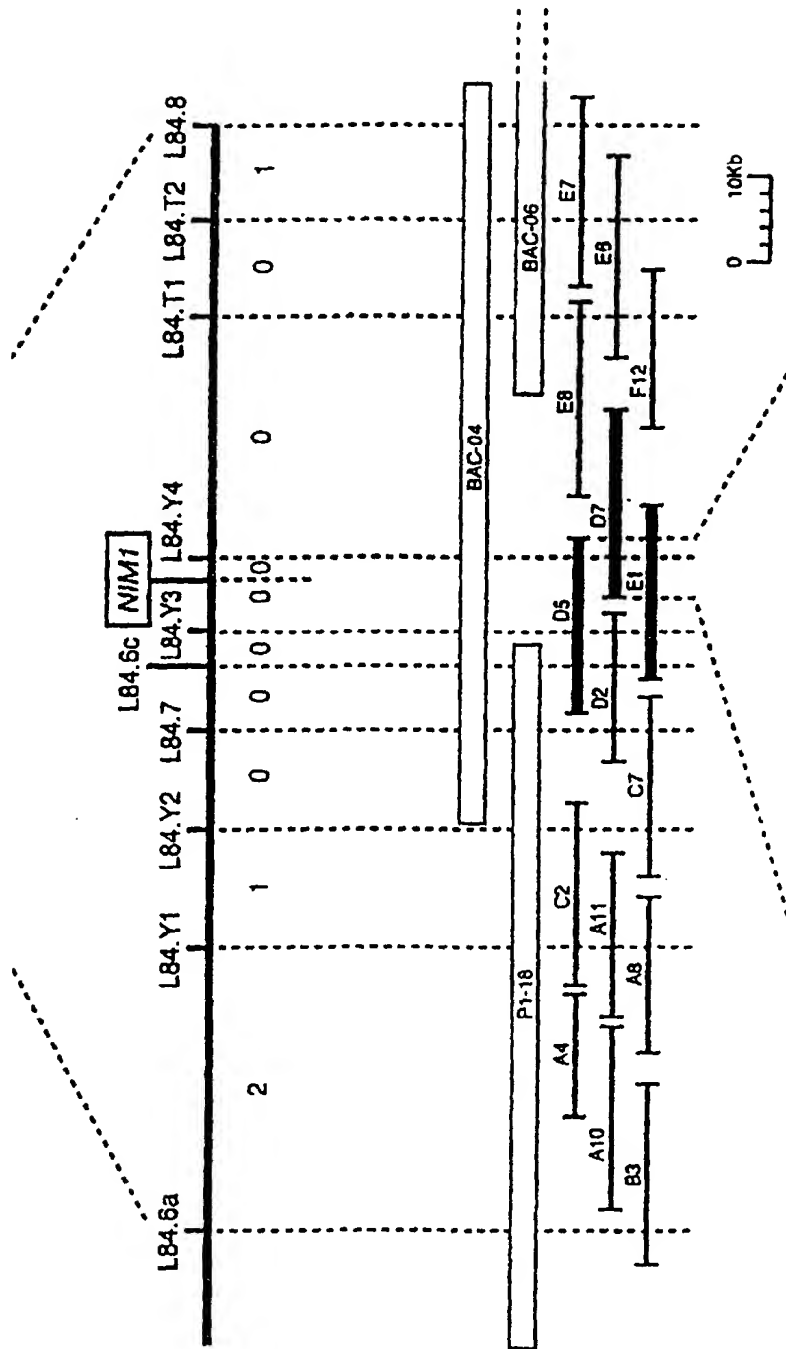
Fig. 12

Schematic representation of the genotypes of the most important recombinants D169 and C105. Shown is the recombinant chromosome with the Ler portion indicated as a black bar and the Ws portion as a gray bar (the non-recombinant chromosome is of the Ws-type). On top the positions and approximate distances of the relevant AFLP markers are shown related to the actual size in kb's as shown at the bottom.

Fig. 13







no. of Recombinants:

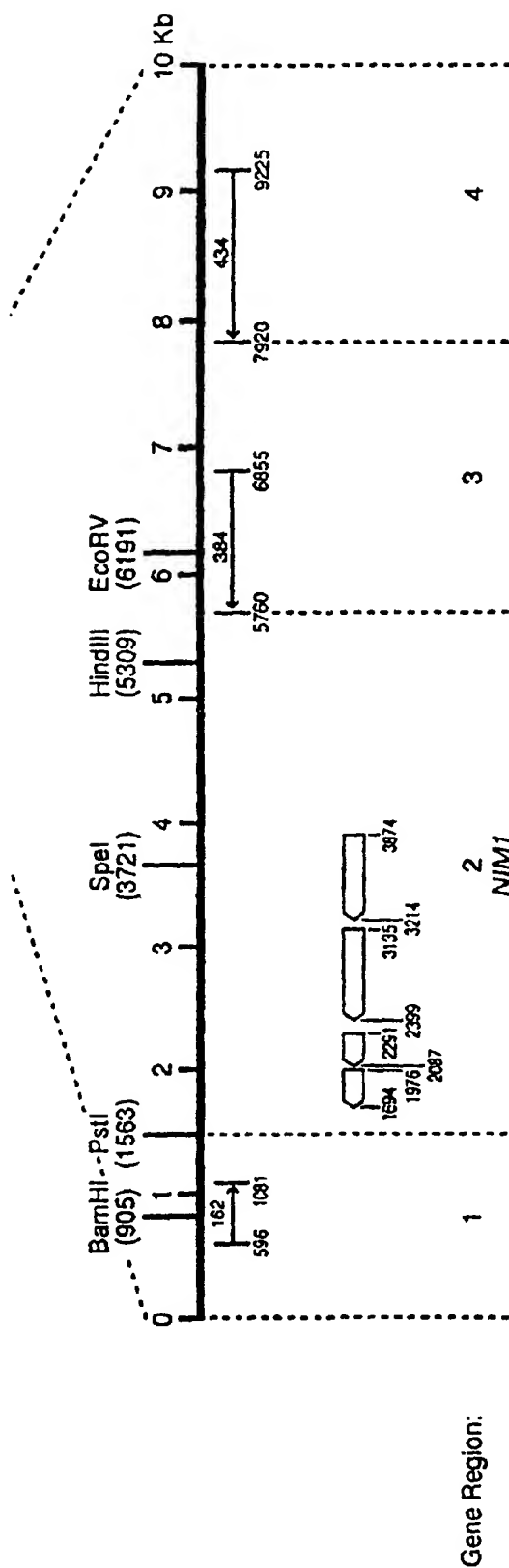


Fig. 14

Nim1 Length: 9919

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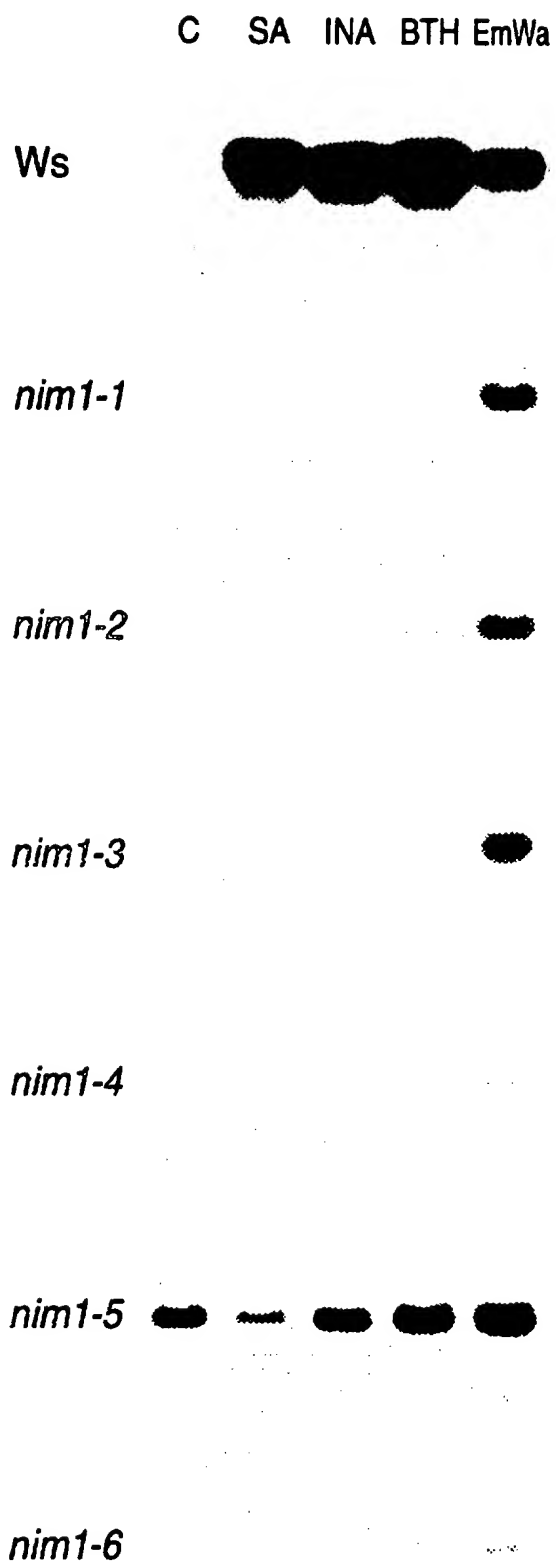
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fig. 17



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fig. 6

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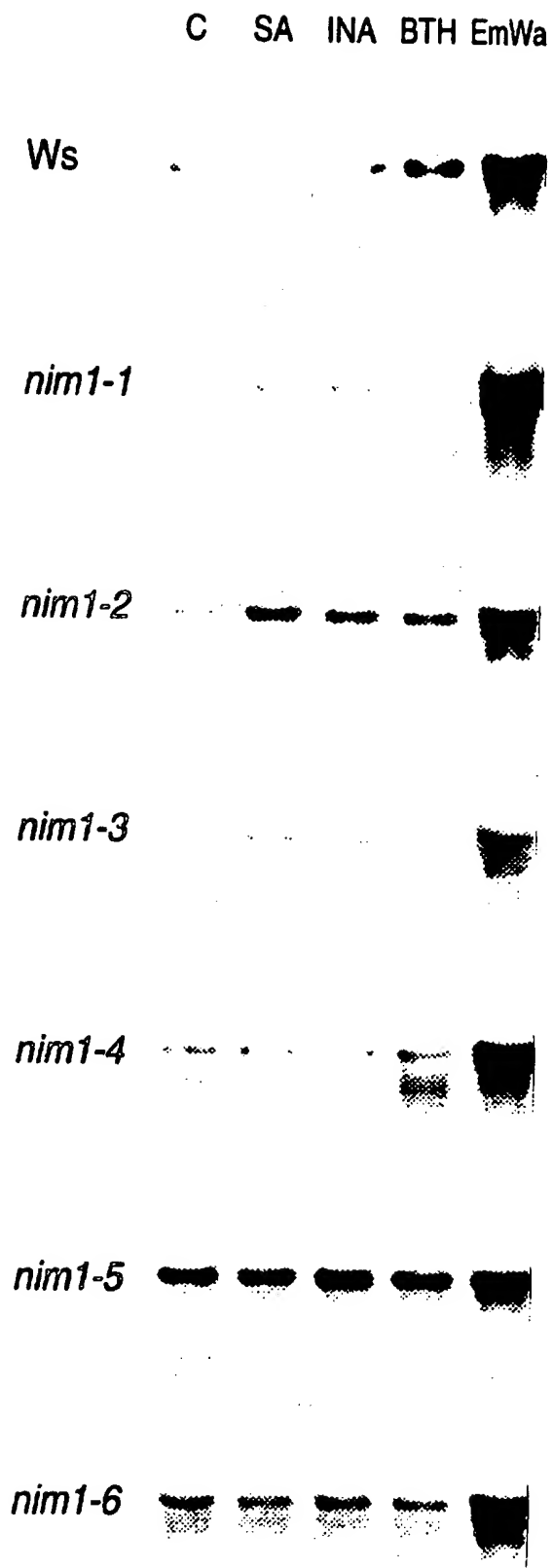
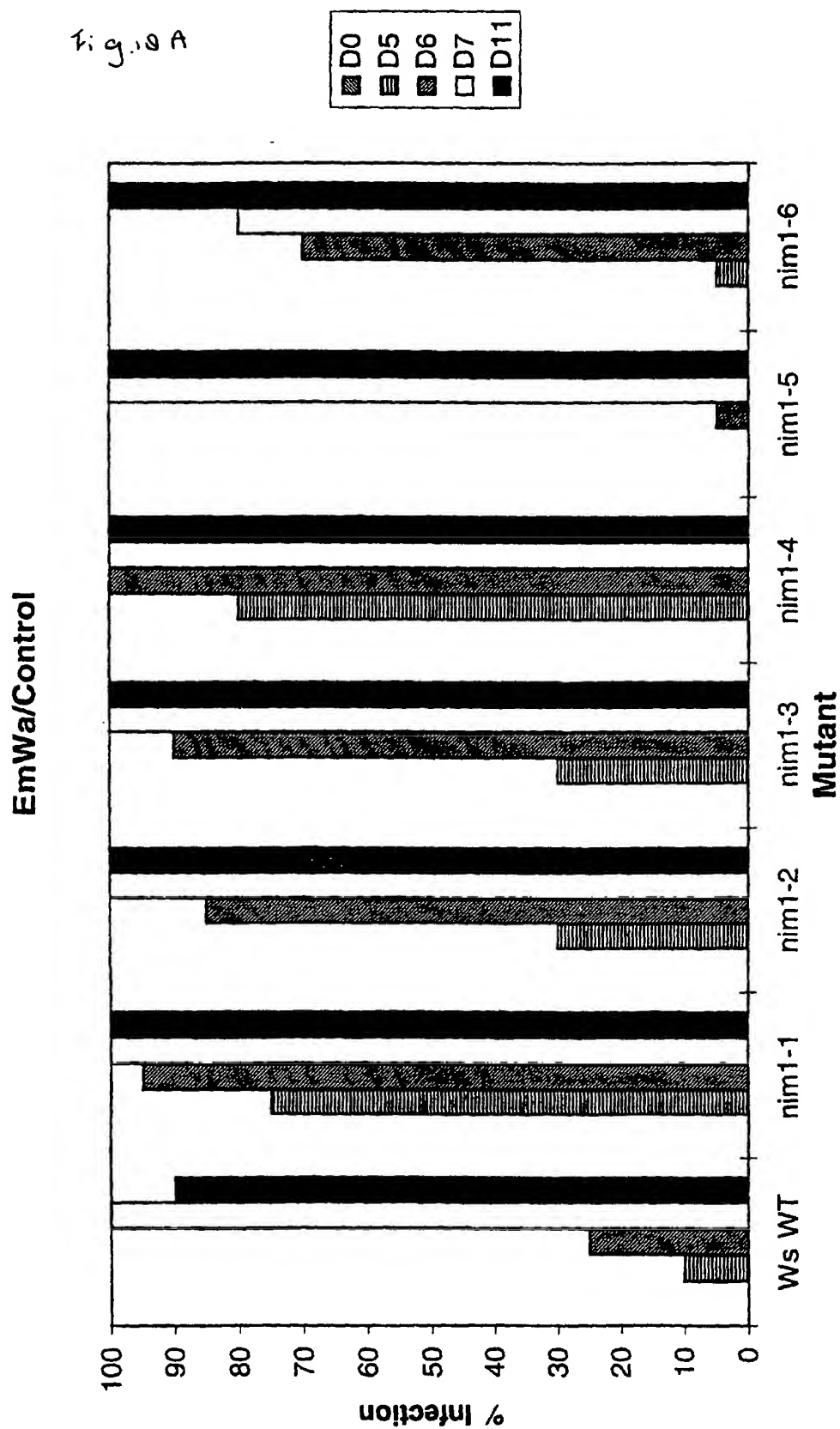
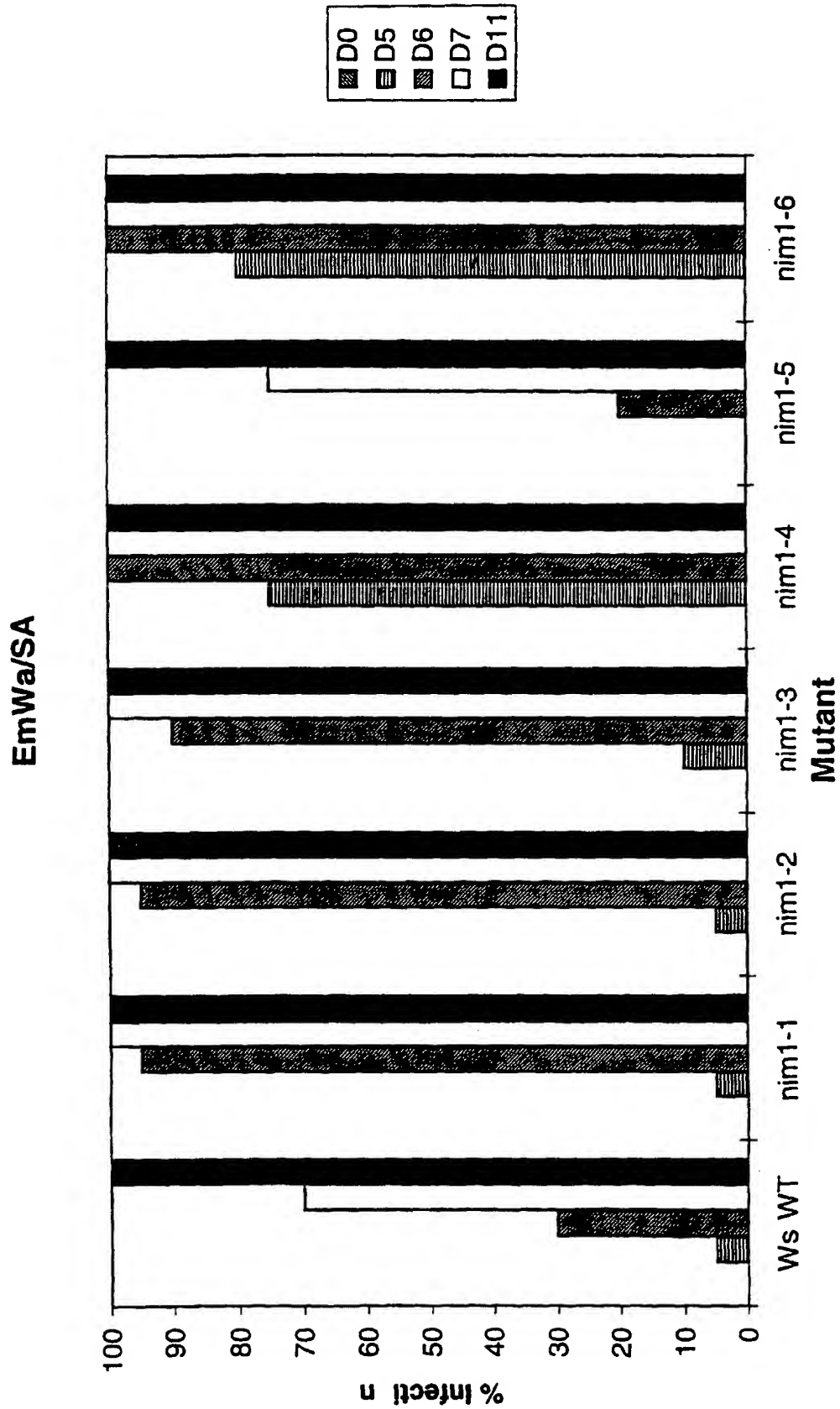
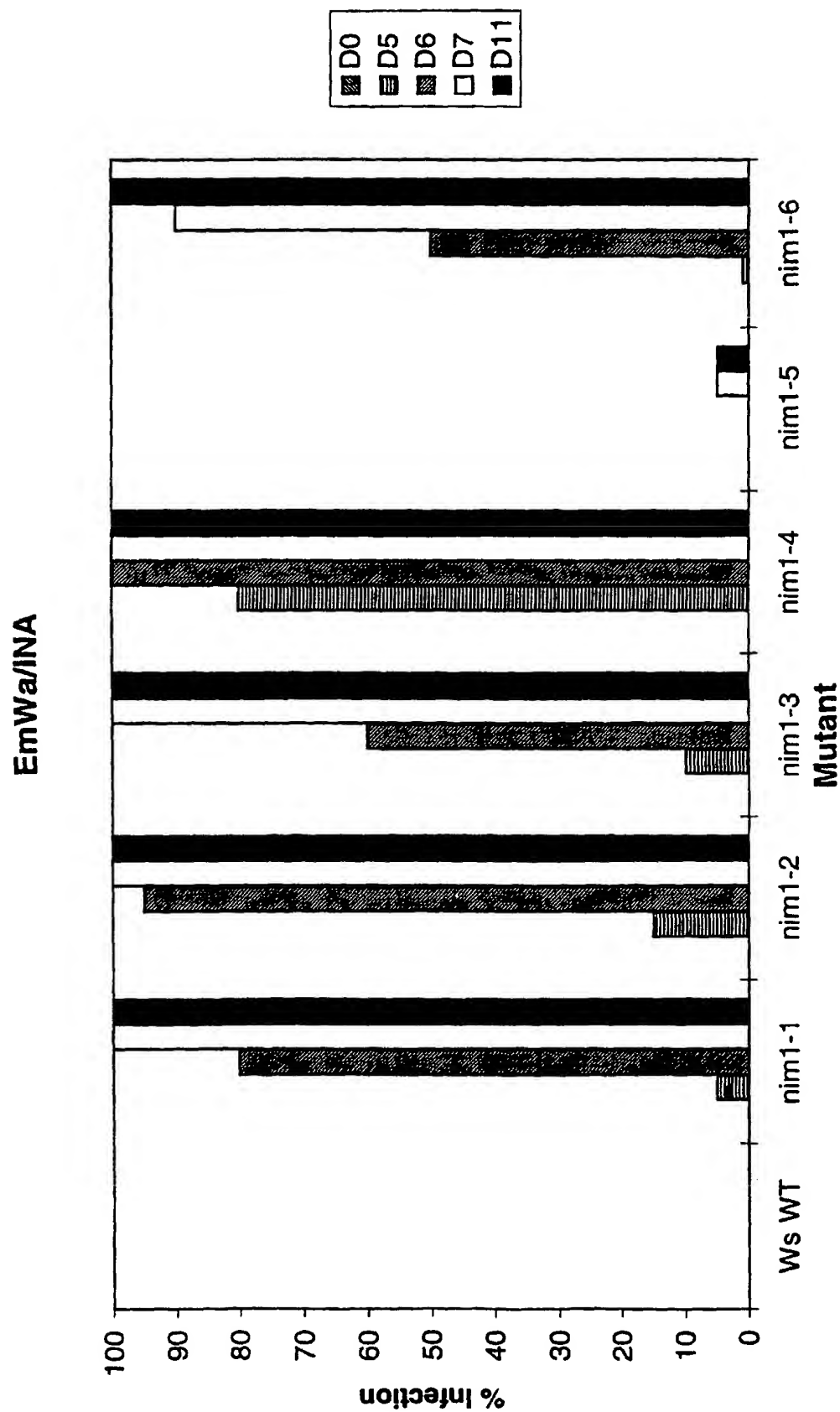


Fig. 9A



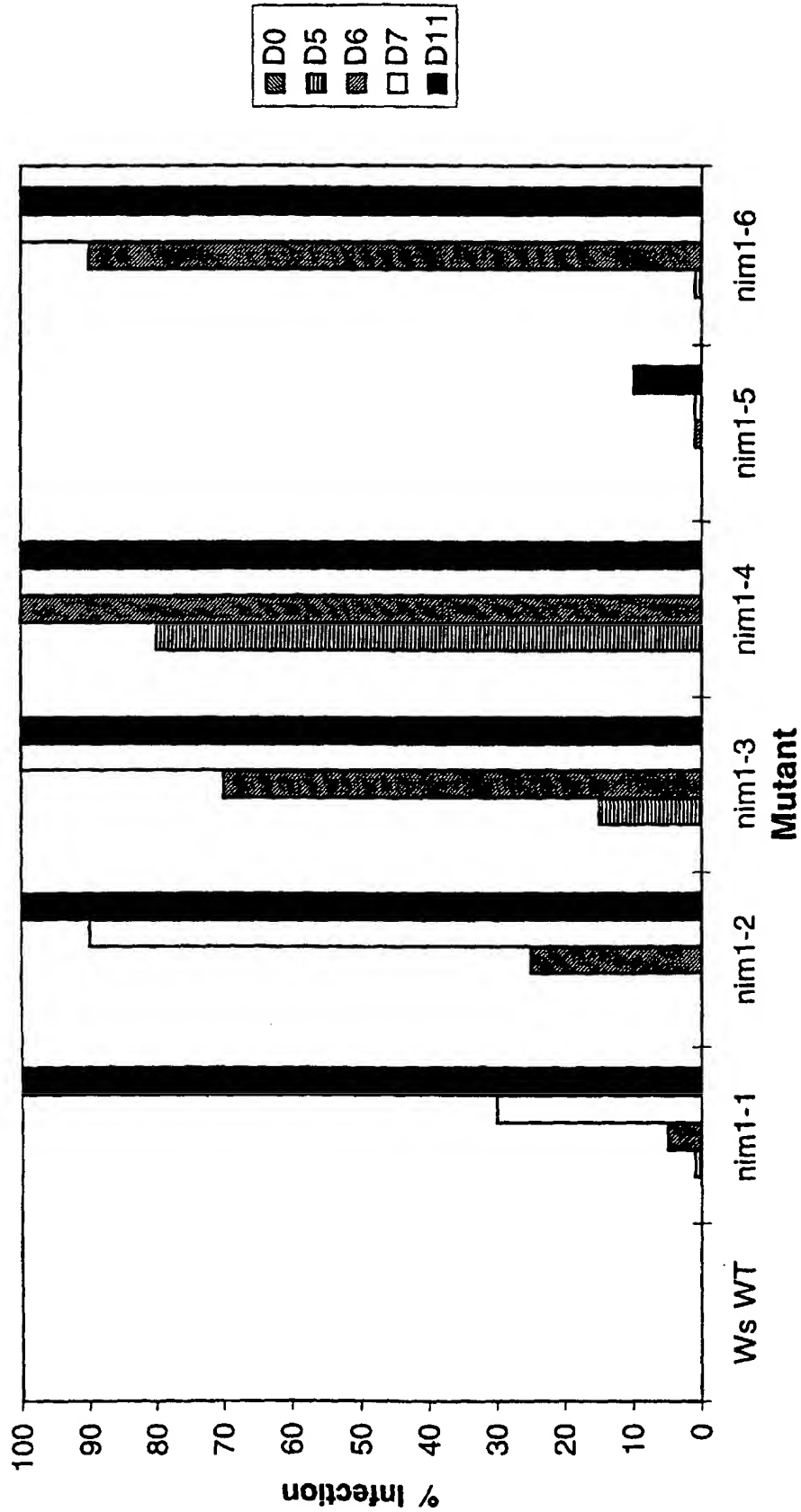
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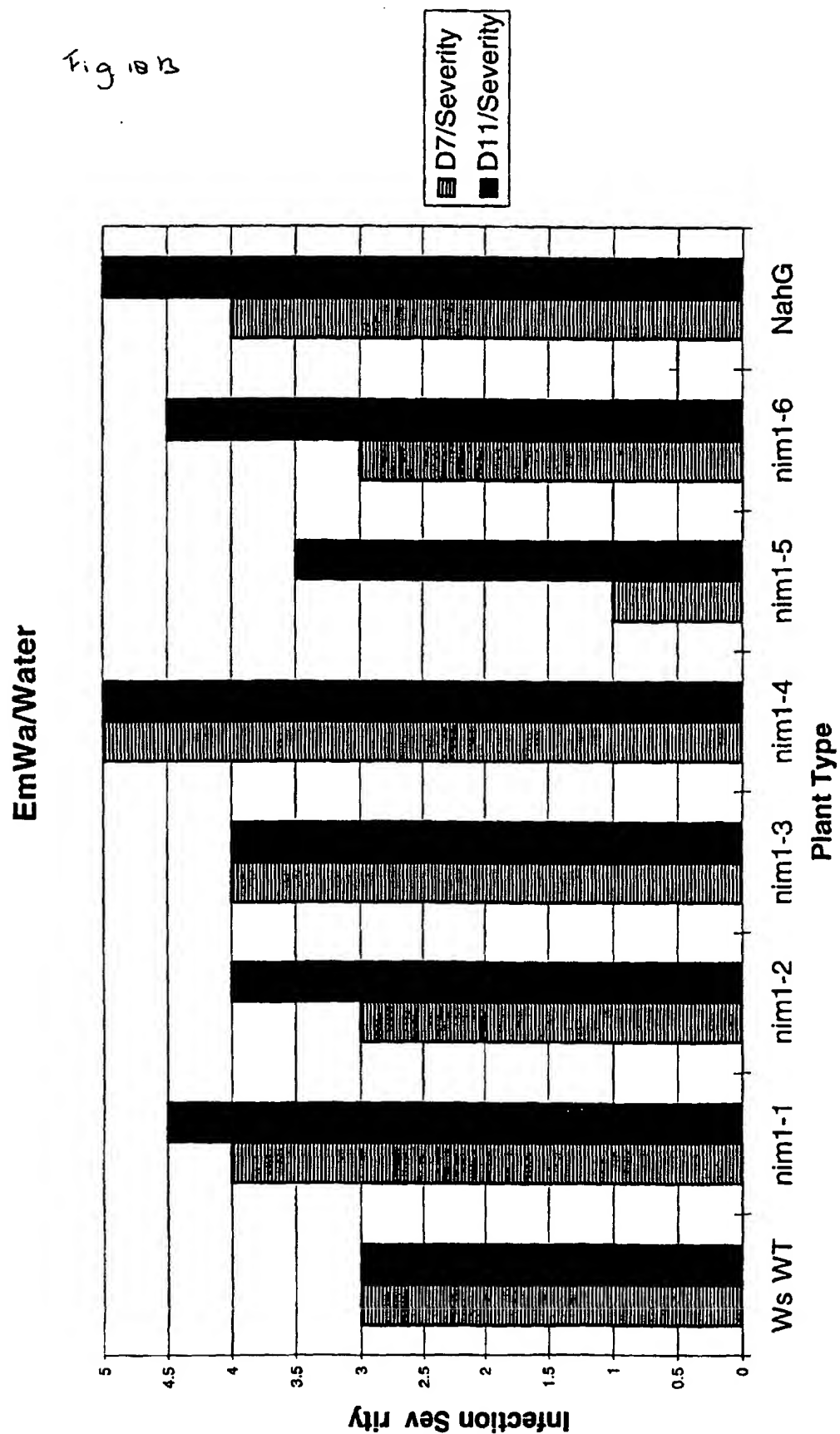
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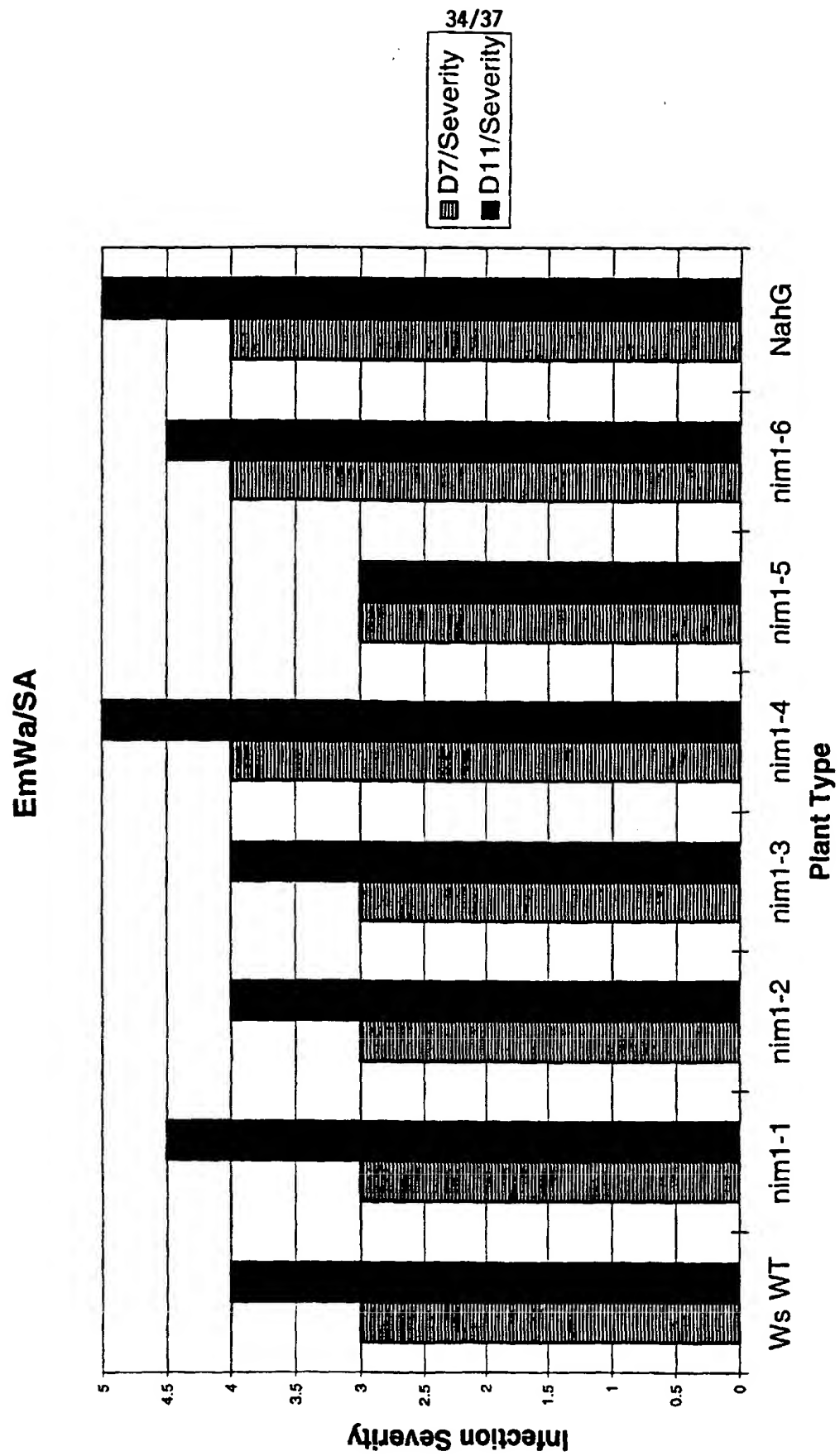
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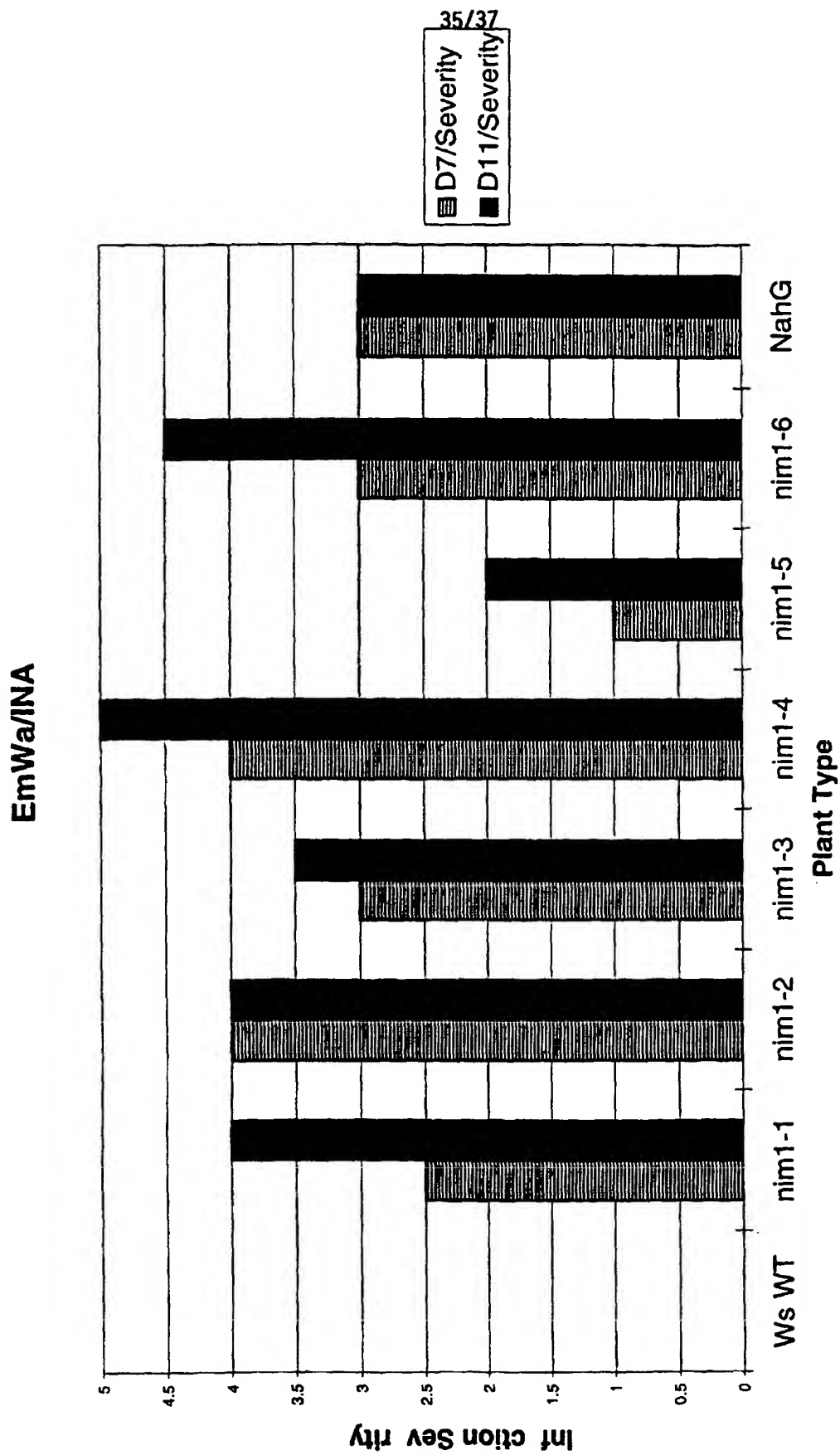


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Fig 10b

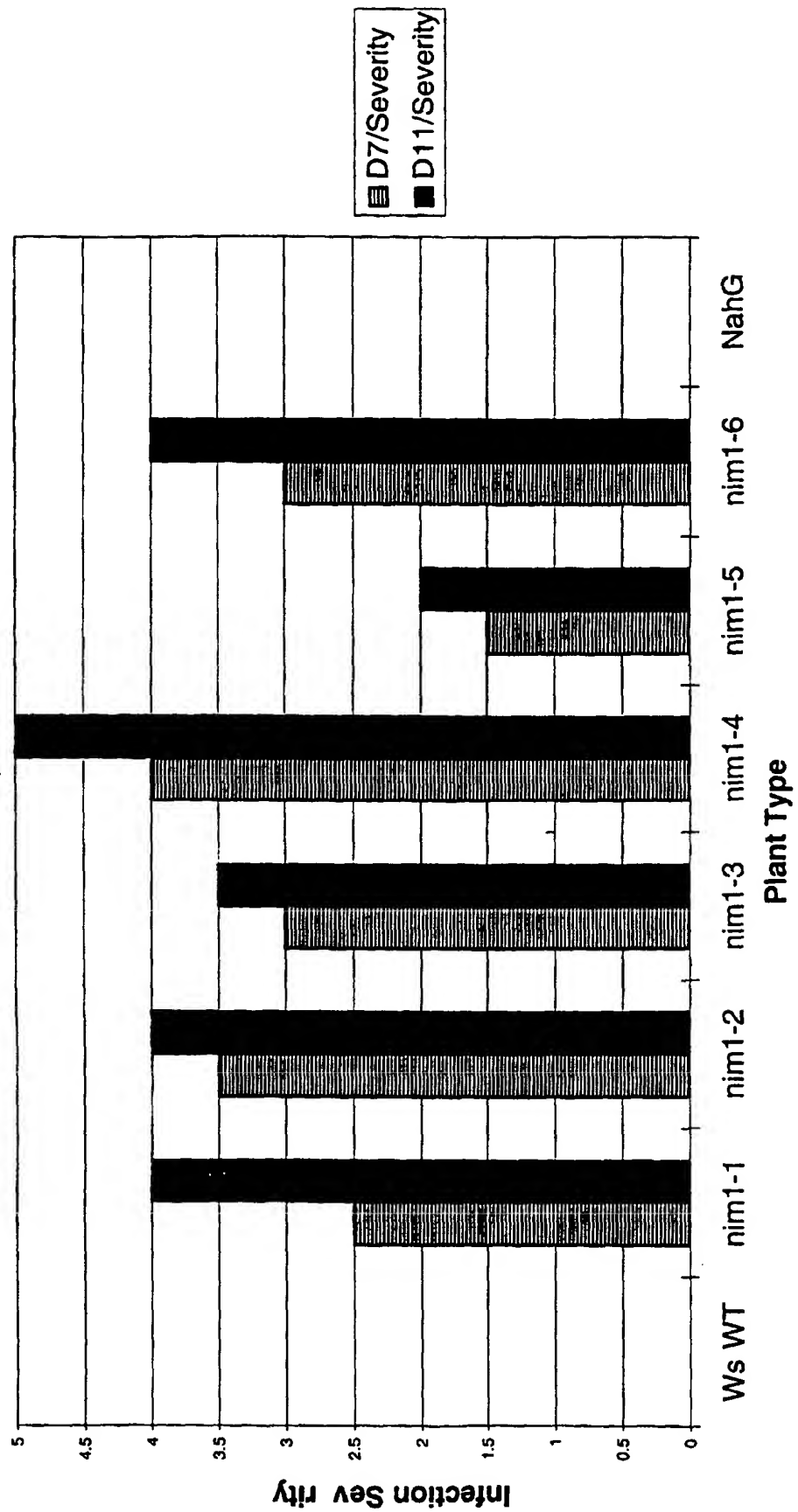






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EmWa/BTH



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nim : 267 VSNVHKALDSDDIELVKLLKEDHTNLDDACALHFAVAYCN 307
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Rice-2 : 33 IRRMRRALDAADIELVKLMVMGEGLDLDLDDALAVHYAVQHCN 155

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nim : 267 VSNVHKALDSDDIELVKLLKEDHTNLDDACALHFAVAYCN 307
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nim : 267 VSNVHKALDSDDIELVKLLKEDHTNLDDACALHFAVAYCN 307
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Rice-4 : 215 PTGKTALHLAAEMVSPDMV 271

INTERNATIONAL SEARCH REPORT

International Application No

PC1/EP 97/01218

A. CLASSIFICATION OF SUBJECT MATTER
 IPC 6 C12N15/82 C12N5/10 C07K14/415 A01H5/00

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC 6 C12N A01H

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

C. DOCUMENTS CONSIDERED TO BE RELEVANT

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X	--- EMBL SEQUENCE DATA LIBRARY, 27 June 1994, HEIDELBERG, GERMANY, XP002034716 NEWMAN, T., ET AL. : "GENES GALORE: A SUMMARY OF METHODS FOR ASSESSING RESULTS FROM LARGE-SCALE PARTIAL SEQUENCING OF ANONYMOUS ARABIDOPSIS cDNA CLONES" accession No. T22612 --- -/-	1,3-5

☒ Further documents are listed in the continuation of box C.

☒ Patent family members are listed in annex.

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Date of the actual completion of the international search

9 July 1997

Date of mailing of the international search report

18.07.97

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Holtorf, S

INTERNATIONAL SEARCH REPORT

International Application No

PCT/EP 97/01218

C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT		
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X	THE PLANT CELL, vol. 6, November 1994, pages 1583-1592, XP002034718 CAO, H., ET AL.: "CHARACTERIZATION OF AN ARABIDOPSIS MUTANT THAT IS NONRESPONSIVE TO INDUCERS OF SYSTEMIC ACQUIRED RESISTANCE" see the whole document ---	1
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